T	able	488	

							00							
	PLACE1009477	13.59	11.11	12.8	10, 3	9, 83	6, 89	8.75	11.15	10.89		- 1		- 1
	PLACE1009493	5.63	4.68	4.85	3.04	3.33	2. 25	3.06	2.29	1.49	**	**	-	-1
	PLACE 1009502	11.43	9.62	8. 39	5. 92	5.81	5. 8	4.63	7.61	3.19		*	-	- 1
5	PLACE1009524	7.56	7. 26	7. 63	1.87	3.86	2.97	2. 66	3, 37	2.45	**	**	-	-1
		18.34	12.36	9.71	6.88	11.49	9.95	6. 88	5.14	3. 52		*		-1
	PLACE1009527							19. 07	17.64	15.03		٠,		1
	PLACE1009531	25. 38	45.76	31.03	26.02	25.63	22.82					**		-
	PLACE1009535	12.3	11.08	8.3	3.83	6.2	5.84	3.36	3.71	4.06	٠	**	-	~
	PLACE1009539	8.83	10.23	7.02	4. 34	7.56	5. 81	5.39	6.61	6. 43		- 1		
	PLACE1009540	37.43	26, 63	27.99	14.5	25.74	28. 63	21	11.9	8. 26		*		- 1
10	PLACE1009542	12	5, 45	5.7	3.62	4, 51	6	3.18	3.69	2. 67		- 1		- 1
	PLACE1009546	12.67	10.42	6.26	3.72	5, 36	5, 98	4.35	10.8	3. 66		- 1		
	PLACE 1009556	7.91	6.37	6.72	5.1	4.62	3.16	3.16	2.46	2.95		**	_	- 1
			15.44	10.67	4,76	13.77	8. 59	5. 94	5. 33	6. 24		**		-1
	PLACE 1009569	11.99							2.83	4.48		**		
	PLACE 1009571	7. 82	9. 37	5. 59	3.8	7.13	3.69	5.84				.		- 1
15	PLACE 1009573	22.09	15.96	12.31	9.76	17.27	9. 7	8. 26	8. 91	7. 87				
	PLACE 1009576	13, 53	9.45	9.65	4.21	6.31	4. 13	5. 49	5.78	4. 15	*	*	-	-
	PLACE 1009580	9.86	9.33	7.56	5.13	7.69	4. 5	8.75	3. 59	4.85		-1		- 1
	PLACE 1009581	12, 95	12.1	8.15	7.19	5, 84	4. 99	3, 45	2.48	3. 53		**	-	-1
	PLACE 1009587	13.3	7.18	8.26	7.06	6.25	5, 41	3.85	4.9	4. 28		1		- 1
		18.08	14. 97	12.54	14.73	10.76	9.75	5. 8	5.66	6.75		**		-1
	PLACE 1009593					14.76	12.77	19. 19	14	13.81		**		- 1
20	PLACE 1009595	24. 28	22.3	15. 19	12.22				2.48	3.3		- 1		- 1
	PLACE1009596	8.31	5. 39	3.71	3.49	5. 52	2. 24	4. 95						
	PLACE1009600	19.52	17.07	12.01	6.59	8.79	3. 99	8.49	7.24	8. 2	. *	*	-	~1.
	PLACE1009604	19.98	10.38	8.96	4. 18	6.3	6. 58	3:83	5.54	4. 55		- 1		- 1
	PLACE1009607	17.2	18	14, 19	8.35	9. 63	7.56	19.54	17.62	14. 33	**	- 1	-	.
	PLACE1009613	8, 31	8, 44	6,06	5.1	6. 18	4. 22	4. 39	2, 55	3. 96		* [		- 1
25	PLACE1009621	18.02	17.88	15.95	7, 12	14.76	8, 85	8.46	6.45	12.83	*		-	- 1
20	PLACE 1009622	16. 93	9, 51	8. 44	5	5. 84	4, 46	4. 96	3, 18	4. 98				- 1
			19.02	15.23	8, 31	9.71	5.05	7. 94	5. 67	7. 85		**	-	- 1
	PLACE 1009624	23.04						3:93	1. 27	4. 29		**		-1
	PLACE 1009637	8. 56	7.93	7.76	3.81	7. 55	4. 55					**		- 1
	PLACE1009639	14.9	6. 92	3. 57	7.62	5. 31	8. 34	5.06	5. 69	7.84		. 1		- 1
	PLACE1009654	23.03	16.88	13.87	7.6	11.88	8.66	7.4	8.09	10.22	*	*	-	-
30	PLACE1009659	17.62	16.71	15. 38	8.59	15.54	12. 83	6.78	8.01	7.95		**		-
	PLACE 1009665	13.55	12, 11	8.08	5.71	10.66	10.87	8. 53	10.59	6.01		1		
	PLACE 1009669	10.91	11.58	11.44	5.71	8. 38	5. 23	8. 32	8.02	8.98	**	**	-	-1
	PLACE1009870	10	4.03	5, 32	3.93	4.76	8, 45	5, 57	4. 18	5. 29			1	- 1
			10. 39	7.63	11.97	12.63	3. 57	6.84	3.75	9, 58				- 1
	PLACE 1009708	13.41						9. 11	3. 24	6. 91			_	- 1
	PLACE1009721	15.79	10, 82	9.31	5. 16	6. 52	4. 42			9. 13	٠.			- 1
35	PLACE1009731	5.6	6. 57	11.06	5.5	8. 58	5. 78	5. 45	4.8					- 1
	PLACE1009735	9.43	10.36	12.52	8, 67	8.77	6.91	8. 15	10.58	12.7				- 1
	PLACE1009737	8.36	8.02	16.98	5.74	17.02	9. 89	4. 98	11.47	4.66			i	
	PLACE1009741	8, 67	7.59	11.34	6.59	7.8	4. 16	2.63	4.89	4.5		*		- [
	PLACE 1009752	9. 51	12.78	18, 39	7.46	12.57	11.15	4.36	9.17	6.85	1			- 1
	PLACE 1009763	16.81	19. 39	15.73	12.5	15, 24	13.69	4.1	8.02	6.41	1	**	l	- 1
40	PLACE1009766	7.54		7.16	6, 98	11. 81	6, 16	9, 17	7. 21	6, 61	1		i i	- 1
		1.34	18.28		16.58		15. 97	7.94	11.57	13.51	ı		ł	- 1
	PLACE1009772	12.62		13.46		26.84		6. 26	4.41	4. 35	1			-1
	PLACE1009782	7.96	6.95	7.99	4. 24	6.2	9. 33				!	•		- 1
	PLACE1009794	8.71	9.98	15. 31	6.91	7.94	5. 64	5.81	8.68	9, 35	١.		ł	- 1
	PLACE1009798	15.7	11.58	19.23	8.55	9. 28	7.81	5.76	14.83	6. 68			- 1	- 1
	PLACE 1009845	5.69	8.07	10.29	5, 79	6.81	9.4	3.27	3.21	4. 35	1	*		-
45	PLACE 1009849			7.54	5, 25	6, 21	5, 82	3.81	4, 56	5. 57				- 1
	PLACE 1009857			9.09	4.47	4. 42	3. 88	4, 36	2.37	3.86			-	- 1
	PLACE 1009861			20. 31	10.08	12. 87	8. 67	13.97	16.84	16, 62	**		-	- i
					405. 22	548.08	335. 22	583. 26	839.73	801.36			1	+1
	PLACE1009872			352.07					34, 28	50.76	**	**	۱.	-1
	PLACE 1009877			121.84	37.51	29.99	31.46	33.83				**		-1
50	PLACE1009879			8.23	4.6	3. 94	4. 45	5.4	6.04	6.14			١-	- 1
SU .	PLACE1009886	6.21	5.71	6.04	3.1	4. 43	3.64	3.76	3.74	3. 91	**	**	1 -	- [
	PLACE 1009888			18.96	7.12	11.3	10.4	7.67	15, 82	6.68			1 -	- 1
	PLACE 1009908			16.94	7.9	15.1	13.2	5.89	21.59	11.16	1		1	- 1
	PLACE 1009919			15. 18	25.14	44, 25	16, 53	6, 38	10.08	17.42			ı	- 1
	PLACE 1009913			13.89	6.9	11. 45	10.84	5.74	7.53	6.61		**	1	- 1
				35.03	28.71	69, 72	45.4	23.4	36.13				1	- 1
55	PLACE 1009923								40, 32				1	- 1
	PLACE 1009924			21.48	6.94	14.99	12.69	29.73					1	- 1
	PLACE1009929	10.23	25.64	14. 3	6.85	14. 35	14. 92	6.32	35.83	7.94	1		1	1

						Table 4	89								
	PLACE1009931			26. 54	12.24	15.5	14. 51	9, 57	9.79	10.08	1 **	**	1 -	-	
	PLACE1009935	8.86	7.44	9.7	2.78	11.02	5.42	14. 95		6.35	1		1		
	PLACE1009947	6.29	6.77	9. 33	4.25	4, 43	3.14	2.04	2, 93	4.01			-	-	
5	PLACE 1009961	7.86	10.21	16.93	5. 24	6.64	4.49	4, 58	6, 45		1		1		
	PLACE1009971	7.79		10.26	3.38	7.18	3. 34	3.67	4. 69	3, 08	1		1	-	
	PLACE1009982	20.35		15, 25	6.6	11, 54	11.1	7.87	5. 59	13. 47	1 .	•	١.	-	
	PLACE1009992	12, 32		11.62	6, 39	8	6.72	5, 24		6. 38	1	i		_	
	PLACE 1009995			27. 33	13.99	19.4	11.61	8, 12	16.7	12.66		i	l _	_	ı
	PLACE1009997	23.76		15.66	6.79	12.9	5.63	10.63	9, 44	8. 19	1	٠	Ι-	-	
10	PLACE 10 10002			12.19	4.66	8.08	5. 35	5. 32	14. 28	7.75			1		
	PLACE 1010011	15. 51		17.86	6, 19	7. 34	6. 29	9.9	6.46	10. 43	**		1		1
	PLACE 1010013			12.68	2.37	2. 99	1. 56	3, 53	4, 24	2. 98	٠٠٠	*	-	~	
	PLACE1010021	15. 95		19.12	7.11	14. 51							1		
	PLACE1010023			21. 32			9.31	5. 22	8.04	6.15	٠.	**	-	-	
	PLACE 1010023	20.87			17.41	29.7	20.44	16.49	7. 28	6.33	ı	*	ı	-	1
15				32.54	10.16	16. 34	7.15	9, 85	8. 36	10.28	•				
	PLACE1010039			10.22	3.99	6.51	3. 25	2.91	6. 23	3.6			ı		
	PLACE1010045		9.8	13.16	5.75	8. 43	6.1	5.11	5.8	5. 51	.*	**	-	-	
	PLACE1010053			9. 04	6.47	4. 19	2.39	2. 32	2. 12	1.16	1	*	ł	-	
	PLACE1010060			15. 21	7.4	8. 59	8.37	12.03	13.21	10.77	**		۱-	-	
	PLACE1010069			13.66	5.33	6. 93	4.79	2.86	4. 42	3.79	•		ĺ	-	
20	PLACE1010070			9.72	2.34	9. 93	4.09	1.65	3. 87	2,87	1	**	l	-	
	PLACE1010074			39.11	24. 21	42.72	26.72	25.32	15. 29	17. 22	1	*	1	-	
	PLACE1010076			55.45	21.22	23.2	19.4	13, 41	12.75	17.59	**	**	۱-	-	
	PLACE1010078			14. 35	6.69	10. 2	8, 63	4, 38	4. 87	4.45		**	-	-	-
	PLACE1010081	27.34	19.16	21, 57	12.55	15.05	11.39	11.73	7, 47	11.77			-	-	
	PLACE1010083	7.1	7.48	5.64	1.97	2.64	2.62	1.69	2.61	1. 96	**	##	-	-	
25	PLACE1010089			9.57	5.5	5. 81	4. 38	4.84	2. 48	4. 43	**	**	-	-	
20	PLACE1010096	14.85		16.35	5, 81	14, 35	9.63	7.28	8, 67	5. 49		•••			
	PLACE1010102	22.29		23. 15	9.26	16.03	13. 51	7.63	9.06	7.61	*	**	-	-	
	PLACE 1010105			15.99	6, 52	12.73	7.6	4.02	4. 82	4.1	*	**	-	-	
	PLACE 1010106			15.73	8, 29	8. 94	12.84	10.96	10.43	10, 44		*	-	-	
	PLACE1010130			4.7	3. 25	3.39	2. 18	3. 26	3, 47	3. 87	:	:	Ι-	-	1
30	PLACE1010132			20.56	20.18	19.1	12, 29	12, 71	11.47	12. 5	١.		1 -		
30	PLACE1010134			9. 66	6.05	6, 05	5, 55	6.72	4, 12	2.88	#	**	_	-	
	PLACE1010139			79, 85	37.26	51, 16	43.94	15.68	26. 17	18.6	::	**	-		
	PLACE1010148		7.43	9. 71	13, 14	7, 54					**	**	-	-	
	PLACE 10 10 155	9.79	7. 58	7.63	3.3	5. 89	4.86 2.96	3. 27 2. 11	4. 13 2. 26	7. 63 2. 53	١.			- 1	
	PLACE 1010156	32. 59	25.06	24.61							1	**	-	-	
35	PLACE1010161	8.48	13.16		14.7	18.09	11.09	7.29	10.58	14.48	٠.	**	-	-	
30	PLACE1010181	8.66	8.08	10.69	5. 15	8. 21	6.03	4. 47	3. 31	5. 46		*		-	
	PLACE 1010 184			6.51	2.62	6.54	5. 02	3.74	4. 12	2. 93		**		-	
	PLACE 10 10 184	8. 57 8. 39	7.46	6.67	3.93	4, 98	3.84	2.86	2.08	2.37	**	**	-	-	
			6.76	12.45	8.31	10.9	5.79	3. 56	3. 42	2.04		*		-	
	PLACE1010231	12.97	10. 31	14. 49	7.96	15.61	7.14	8. 93	9. 42	8. 37		*		-	
40	PLACE1010235		15	11.24	4.59	11.54	4. 53	6.11	5.14	4.76		**		-	
40	PLACE1010237	5.04	3.77	4.4	2.18	2. 37	2.25	2.72	0.64	1. 32	**	*	-	- [	
	PLACE1010251	9, 46	4.47	6.29	5.09	4.79	10.14	3.78	5. 92	3.88					
	PLACE 1010261	8.26	6.41	4.7	3.42	2.86	2.61	4. 52	5.73	4.64	*		-	- 1	
	PLACE1010270	7. 53	8.07	6.36	3,29	6.44	3. 19	3.88	4. 95	3. 8		**		-1	-
	PLACE 1010273	8. 97	10. 55	5. 7	2.72	5. 68	3.04	2.78	2. 04	3. 83		*		-1	
	PLACE 1010274	20.97	18.72	29, 56	10.88	16. 29	14.38	6.73	5.73	6.16		**		- 1	
45	PLACE 1010277	15.29	14.99	14.3	8.87	13, 93	8, 53	11. 24	4.25	6,68		*		- 1	
	PLACE1010293	16.94	20.61	17.07	9.41	11.63	8. 24	17.5	8.1	13.23	**		-	- 1	
	PLACE1010297	38.08	33. 94	34, 95	17.91	31.69	15.63	31.8	20.1	25.32				- 1	
	PLACE 1010300	21.55	15.58	11, 87	9.15	8.78	7.88	3, 82	5, 36	5.08		*		- 1	
	PLACE1010310		293. 14	231. 29	170.36	221.96	136.69	214, 56	235, 35	207. 22	*		-	ı	
	PLACE 1010321	10.7	10.58	7.81	4. 96	7, 93	6, 91	5.6	6.6	6. 56				- 1	
50	PLACE 1010324	6, 25	5.69	6,54	2.88	4, 51	2. 6	4, 43	3. 34	2.72		**	_	-1	
	PLACE1010329	14, 25	11.37	11.93	5. 93	10.64	4, 89	9, 64	9. 84	9, 32	Ĭ	7		_	
	PLACE 1010330	12. 28	14. 21	10.49	11.84	10.18	7.63	12	7.94	17.86		1			
	PLACE 1010335	27.7	52.66	38. 23	18.39	9.7	13.5	16.61	15.38	28.61			_	- 1	
	PLACE 1010341	6.44	4. 91	4.72	2	3.7	1. 35	4, 14	2. 93	4. 21	•		-	- 1	
	PLACE 1010342	2. 35	2. 93	2. 85	2.16	1,61	1.84	1.14	1.19	0.83	:	**	Ĩ.	_ !	
55	PLACE1010346		6.88	7. 23	4.08	5.74	2.77	5. 84	6, 92	5. 88	:	**	Ī	1	
	PLACE1010362	13. 25	8. 57	9. 24		7.05	3, 39	11.5		14.05	*		-	_ 1	
	B	14.75	U. U!	3. 44	U. 98	1.00	3. 33	11.0	13.76	19. 05					

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						Table 4	90								
	IPLACE1010364	11.29	7.43	7.221	2, 26	3.51	2.16	4. 95	3, 65	4, 16		*	-	- 1	
	PLACE 1010368	10. 78	8, 61	7. 03	3.98	6, 56	5, 19	7.58	7.15	5, 73				- 1	
	PLACE 1010373	30. 2	28, 61	24, 17	14.82	22.58	10, 46	24.63	20.09	20.87	*		-	- 1	
5	PLACE 10 10383	18.56	18, 15	16.04	10	15, 54	3.52	27.13	12.41	12.03				- 1	
	PLACE 1010385	2.48	1.73	1.18	- 0.83	2.33	0.49	2.65	1. 19	2.06				- 1	
	PLACE1010389	8, 48	7.51	8, 42	3.75	4, 47	5. 47	3. 96	4. 51	2. 91	**	**	- 1	- 1	
	PLACE 1010401	3. 24	0.96	4.6	2.29	2, 49	1.63	2. 81	2.31	1.6				- 1	
	PLACE 10 10410	22.91	15, 39	21.28	8, 31	20, 95	12.79	12,79	10.9	11.3				-1	
	PLACE 1010418	18.78	19.01	17, 17	5.75	13, 45	6.67	11.25	18.37	11, 11	*	- 1	-	- 1	
10	PLACE 1010425	10.58	10, 75	18, 36	4. 2	4. 88	3, 19	8.87	11.17	8.75			- 1	- 1	
	PLACE1010443	12.48	14, 66	16,77	5.6	14, 99	6.09	8.79	8. 23	11.76			1	-	
	PLACE1010445	36, 17	45.7	37, 43	17, 56	20, 28	18.29	39. 23	32, 46	41, 56	**		-	- 1	
	PLACE 1010481	60.4	46, 95	65, 37	22.92	26, 42	21, 44	27, 71	30, 53	27.4	**	**	-	-1	
	PLACE 1010482	127.4	82.31	84, 53	104, 64	120, 51	109.85	62.48	50.87	49. 97			1	-1	
	PLACE 1010491	71.28	72.3	73, 41	43.51	47. 95	35. 09	68.63	86.51	74, 84	**	- 1	-	- 1	
15	PLACE1010492	20. 32	20.74	19,72	20,62	27. 04	23. 27	9.07	12.55	10, 15		**	ı	- 1	
	PLACE 1010509	11.2	13.07	17.83	8, 18	12, 58	8, 24	5.79	8, 34	7.68		*	1	-1	
	PLACE 1010518	9, 76	12.31	18,67	8, 17	8, 57	7. 5	8.04	6.33	6, 73			1	- 1	
	PLACE1010522	8. 32	9, 35	9.63	6, 71	11, 42	6. 63	6. 37	5. 88	7.65		*	l	-1	
	PLACE1010529	13.09	22, 15	14.64	11.31	17.05	8, 58	7.61	9.45	9, 29			i	- 1	
20	PLACE 1010547	36.79	34.33	38, 34	12.39	16, 81	11, 19	21.21	24.3	23.05	**	**	i -	- 1	
20	PLACE 1010560	10, 15	9, 34	9, 56	6.09	4, 54	5, 89	6.02	5.35	6.13	**	**	i -	-1	
	PLACE1010562	4,79	4.39	12.01	8.8	8.65	6.38	6.41	7.48	4.4			i	- 1	
	PLACE 1010579	74.54	67, 98	59.08	46, 33	48, 92	48, 24	56. 27	84. 49	60.31	*		-	- 1	•
	PLACE 1010580	38, 79	45.7	46.75	22.24	30	17, 17	31.95	38. 27	36. 2	*		-	- 1	
	PLACE 1010599	29.35	25, 54	32, 71	12.61	15, 12	14, 18	19, 83	24. 34	23, 16	**		-	- 1	
25	PLACE1010606	31.76	30, 37	40.62	15.46	19. 15	14.23	22.77	25.83	28.81	**		-	- 1	
	PLACE1010616	16.39	26, 11	18.04	12.86	17.78	7.33	3.49	3.67	4. 92		**	i	- 1	
	PLACE1010622	9.08	4.75	12.08	2.52	4. 26	2.63	11.54	11.12	6.28			1	- 1	
	PLACE 1010624			204.46	212.8		210.51	166: 11	171.99	155.01			l	-1	
	PLACE 1010628	48, 65	44.83	51.05	27. 25	26, 82	37,41	15.79	19.04	21	*	**	i -	-	
	PLACE1010629	19.03	17.43	20.71	12.12	16.38	15.36	14.93	10.66	11.67	*	*	-	- 1	
30	PLACE 1010630	15.62	16.61	29.46	8.82	19.7	10.88	11.96	14.1	11.42	1		i	- 1	
	PLACE1010631	22. 99	34. 46	25.54	20.07	21.82	17.69	19.23	10.89	16.98			i	- [	
	PLACE1010651	8		11.34	5.4	9, 54	5.75	4.35	5. 43	5.06		**	ı	-1	
	PLACE 1010661	13.76	16.34	32.78	7.67	13.64	27.31	6. 28	10.55	7. 26			į .	- 1	
	PLACE1010662			30.53	8.67	12. 27	9.85	25.45	28.04	24.56	**		-	- 1	
0.5	PLACE 1010668	37.33	37.7	40.55	22.62	25.66	20.72	36.78	44. 21	37.45	**		-	- 1	
35	PLACE1010702	12.3	6. 26	13.51	6, 5	7.73	5.48	8. 16	17.08	6. 46	١		1	-1	
	PLACE 1010709			68.95	55.09	47,9	57.04	50.07	52.69	55. 39	**	**	-		
	PLACE1010713	80. 41	68. 48	77.71	36.46	48. 82	48. 26	28. 63 3. 33	32.14 8.97	23.64 5.69	**	**	-	- 1	
	PLACE 10 10714			15. 24 17. 8	4.19	7. 98	4. 28	5.99	13.73	3.7	**	•	-	- 1	
	PLACE1010716					22. 94	9.1			3. 55				- 1	
40	PLACE1010717			15. 64 34, 2	13.63 14.65	18. 64 18. 34	10. 49 8. 25		15. 16 22. 85	22.74	**	*	1_	_ 1	
40	PLACE1010720 PLACE1010739	32.05		24.71	15.33	21.42	15.16	17.5	28. 68	18.95	*	٠	ΙĪ	- 1	
	PLACE1010743	44.76		34.77	15.52	24.11	19, 41	14.02	9. 67	5.8	1:	**	-	_ 1	
	PLACE1010752	26. 13	19.45	22.02	11.29	9, 49	11.37		24. 35	13. 97	**	••	-	-1	
	PLACE 1010751	15, 13		32. 55	15.98	9. 45	8.78	11.14	13.99	12.89	1 **			- 1	
	PLACE 1010771			37. 99	25.83	90, 51	39. 2	31.02	18.86	25. 47	l			ı	
45	PLACE1010784		116, 91	93. 11	48.56	63, 14	52.26	30.32	31.37	33.38	**	**	-	- 1	
	PLACE1010786	16. 19		16. 87	8.23		9. 85		6. 49	9. 87	1	**	l	-1	
	PLACE 1010789			40.6	14.8	19, 17	10.09		47.22	35, 88	**	•••	-	- 1	
	PLACE 1010800			14.67	8.91	6.6	5. 82		5.43	4.76		**	۱-	-1	
	PLACE1010802			27. 18	16.21	23. 96	4, 91	12.32	5.78	37.85	1		1	1	
	PLACE 1010811	26.9		31, 28	13.09	18.01	12. 69	11.33	11.86	10. 25		**	-	-1	
50	PLACE 10 10813			9.67	4.84		4. 87		2.83	2.64		**	-	- 1	
	PLACE 1010827			12.05			5, 87		5.02	3.76		**	-	- 1	
	PLACE 1010833	9. 18		8.82	3. 14				2.07	1. 97		**	l -	- 1	
	PLACE1010839			20. 18	11.7	20			11.13	15.42	1	*	1	-1	
	PLACE 1010858	15.47		17.88	6.16		4. 26		8. 32	12. 25			1 -	1	
55	PLACE 1010857			18.71	21.99				15. 82	15, 61	1		1	- 1	
30	PLACE1010870			4.28	3.87				2.04	1.94		**	l -	-1	
	PLACE1010877		20.6	20.89	14.13	8. 42	9.71		12.71	7.25	**	*	l -	-	

					1	able 4	91								
	IPLACE1010882	17.06	13.54	8. 421	6.17	5.69	5, 82	7.13	5.03	5. 37	*	- 1	-	-	
	PLACE 1010891	27.43	30, 32	21, 15	13.46	14.41	7.46	15.38	15.07	24, 71	*	- 1	-	- 1	
	PLACE 10 10896	13.62	18.23	16.16	6. 92	8, 03	5. 78	3, 54	4.62	4.06	**	**	-	- 1	
5	PLACE 10 10 900	66.6	75.65	67, 62	55.68	59.03	45, 47	67.12	53.92	39, 14			-	1	
						35.16	33.1	29.84	23.73	27. 17	**	**	_	-1	
	PLACE1010916		71.53	76.5	35.13									-1	
	PLACE 1010917	14.21	14.67	11.27	7.82	5. 59	7.41	7.69	7.06	8.76	**	**	-	~ [	
	PLACE 1010924	5. 86	2.7	4. 5	2.89	4.47	3.08	2.47	3. 1	1.31		- 1		- 1	
	PLACE 10 10925	4.94	4, 55	4.99	1.88	2. 53	1.67	3.09	1.69	1.78	**	**	-	- [	
	PLACE 1010926	13, 94	11.16	11.63	5. 87	7.64	6. 22	5. 6	4. 23	4. 29	**	**	-	- 1	
10		207.77				130.24	102. 36	197.42	120.16	123.73		- 1	-	- 1	
	PLACE1010943	31.5	25. 2	22.98	12.49	22. 16	9	15, 12	16.22	13, 62		* 1		- I	
		83.2		68, 43	51.82	78.63	47.87	87	58, 47	73, 16		. 1			
	PLACE1010944		92.75					22.66	11, 99	13.89		- 1	_	- 1	
	PLACE1010947	33.76	22.26	22. 3	12.84	13.89	8. 22					1	-	- 1	
	PLACE1010954	10.33	11.3	11.05	5. 3	5.81	5. 51	5. 34	5.12	4. 22	**	**		- 1	
15	PLACE 1010960	15, 62	30, 44	20.31	7.94	8.28	7.04	11.37	9, 05	8.5	*	*		- [	
	PLACE 1010965	13. 17	14.34	9.79	7.54	4.78	3.78	4. 15	4.73	5. 18	*	**	-	-1	
	PLACE 10 10968	18, 41	14.95	14, 27	6.84	9.6	5. 59	12,62	6.34	9.37	**	*-	-	- 1	
	PLACE 10 10978	57.65	54, 74	36.05	40.68	34.09	13.7	16, 85	10.36	20.32		*		-1	
	PLACE 1010982	21.44	26.06	15. 9	7.42	6.54	11.09	10.66	7.85	9. 32		*	-	-1	
	PLACE 10 10 990	327. 98	344.55				240.66	208.45		173.44		* 1		-1	
		100. 16	59. 4	54. 99	23.72	27. 4	15.8	91.94	75.46	99.74		٠ ١	_	- 1	
20	PLACE 1011017								7.41	7. 81	•	- 1		- 1	
	PLACE 1011019	13.37	8.64	6.46	8. 37	6.37	4. 79	6. 61				- 1		- 1	
	PLACE 1011026	17.67	16.91	13.69	8.74	11.48	8. 93	15, 71	13.37	15. 94	. *		-		_
	PLACE1011032	5. 5	8.88	7.51	2.84	5.75	2. 99	4.5	3. 38	3.21		*		-1	
	PLACE1011041	11.32	11.72	11.41	4.79	6. 17	4. 9	2.74	0	1.85	**	**	-	-1	
	PLACE 1011045	30.68	27.07	24,09	20.73	22, 77	14. 24	25, 88	22. 96	30.06		- 1		- 1	
25	PLACE 1011046	10,75	10.03	9,15	3.68	5.42	4, 38	4. 21	3.5	2.56	**	**	-	- 1	
	PLACE 1011054	23, 11	26.67	18.54	13.24	17.45	11.68	27.3	21.77	27. 93		- 1	-	- 1	
	PLACE 10 11056	5. 67	3, 31	3. 16	3.73	4, 11	8. 46	6.33	3.05	6, 91		- 1		ŀ	
	PLACE 1011057	10.04		7. 99	8, 15	6. 14	3. 72	11.95	6.55	7.6		ı		- 1	
					7.08	9, 45	9. 52	6, 53	6.86	9, 19		. 1	_	. 1	
	PLACE 10 1 1059	19.4		11.33						0.49				-1	
	PLACE1011066	9.28		5. 97	5.75	7.03	4.1	2, 13	5. 18			٠,۱		- 1	
30	PLACE1011087	26.84		33.04	57.32	27. 46	16. 29	127.9	19.42	37.4				- 1	
	PLACE1011090	10. 52		6.48	3.01	5. 18	3. 1	6.04	4.61	6. 16	*	- 1	-	- 1	
	PLACE1011109	43.09		13.76	6.5	19.94	4. 84	12.2	7. 21	13. 97		1		- 1	
	PLACE1011114	17. 98	24.04	19.89	13.12	19, 15	8. 13	12.09	13, 25	14. 96		*		-1	
	PLACE1011116	30.03		37.37	10.98	8.87	7	8. 26	10.18	5, 99	**	**	-	- 1	
	PLACE1011122	5.5		5.93	4,16	4, 82	3.11	3.97	4.69	2, 89	*		-	-1	
35	PLACE1011133	6.47		6.51	3.81	3, 93	3. 81	2.72	5.84	2, 91				- 1	
		5.48		6. 34	4. 28	4. 27	3. 64	3.72	3.86	3.76	**	**	_	- 1	
	PLACE 1011134				2.21	5, 45	3. 27	2. 93	3. 83	4, 13				- 1	
	PLACE1011143	4.75		5. 59					6, 69	4. 48		**		- 1	
	PLACE1011146	10.48		12.48	9.3	11.11	4.11	4. 38				**		- 1	
	PLACE1011160			6.71	4.98	1.77	3. 25	6,45	4. 47	3.78					
	PLACE1011165	21.99		32.58	8.87	19.75	11.93	12.6	15.59	9. 42		*	-	-1	
40	PLACE 1011181	5. 25	3.37	6	2.06	4. 49	2. 29	2.61	4.09	3.85				- 1	
	PLACE 1011185	60.3	35.78	36.17	37.68	34. 15	24. 31	17. 32	33.87	29.05				- 1	
	PLACE 1011186	25.07				20. 81	15, 83	19, 93	27.9	21.4	*		-	- 1	
	PLACE 1011203					12.5	6, 63	8.02	13, 49	10.02				- 1	
	PLACE 1011214					5. 78	3. 95	1, 49	2.02	3.84		**	~	- 1	
	PLACE 1011219					4. 52	4, 58	9.07	4.64	3.23	ı .			. 1	
45						10. 51	4, 92	4, 92	5. 52		1	**	1	-1	
	PLACE1011221							5, 43	7.1	4. 99		**	١_	-1	
	PLACE1011225					7.48	4, 86				١	**	_	_	
	PLACE 1011231					24.8	13, 83	19.12	24. 97	26. 97	١			- 1	
	PLACE1011236	13.0			5.99	7. 59	5, 06	4. 42	6.98		**	**	1-	-1	
	PLACE1011247	571.8	409, 73	639.71	528.63	738.01	566. 23	553.23	542.58	267.58	l		l	- 1	
	PLACE 1011261			9.65	4.09	5, 29	4. 39	2.77	4, 83	3. 37	1		•		
50	PLACE 1011273					7, 81	4, 87						-	-1	
	PLACE 1011278					19, 17	11.18		5. 1		1	**	1	- 1	
	PLACE 101128					7. 47	4. 38				۱.	**	- 1	_ 1	
													i	- 1	
	PLACE1011291					1. 37	0.09				1 **		١.	1	
	PLACE 1011291					6.09	7. 28						1		
55	PLACE 1011310					24. 65		26.25				*	1 -	-	
	PLACE1011311			13, 64								**	1 -	- 1	
	PLACE 1011321	316.1	1 225, 18	251.29	230.23	241.81	208, 08	340.56	284, 43	231.07	1		1	- 1	

PLACEIONIZES 67.48 86.77 34.36 32.79 55.55 25.52 25.00 8 16.56 23.25 8 8 9 -   PLACEIONIZES 77.6 32.89 20.69 24.89 22.44 17.06 26.8 25.99 33.6 8 3.83 10.76 1 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9							Table 4	92			•				
PLICEIDITISAD   6,86   4,22   8,9   7,42   3,33   10,95   4,69   3,122   3,76		PLACE1011325	67.49	68. 67	34.36	32.79	55. 55	25. 52	26.06	16. 66	23. 25		*		- I
PLACEIDISS   1,06   9,45   9,31   4,05   5,07   3,21   9,83   8,81   10,17   17   18   19		PLACE1011332	37.6		20.69	24.89		17.06	26.8		39. 36		- 1		-
PLACEIDISSO 7.09 6.29 7.09 5.01 4.54 2.00 4.25 9.22 7.13													- 1		- 1
PLICEION 364 36.2 48 46.89 18.22 21.7 16.48 29.76 22.08 21.29 \$ \$ 4   PLICEION 377 86.47 40,06 44,55 30.4 25.77 22.53 35.29 36.44 48.61 \$   PLICEION 377 86.47 40,06 44,55 30.4 25.77 22.53 35.78 24.61 \$   PLICEION 377 9.70 4 6.27 10.93 2.06 3.17 16.52 2 PLICEION 378 18.03 11.85 15.3 3.37 4.12 3.15 3.67 7.38 4.18 \$   PLICEION 378 18.03 18.05 19.39 2.06 3.1 2.3 1 1.2 3.77 5.3 8.4 18 \$   PLICEION 378 18.30 14.91 27.77 5.31 8.25 18.52 18.52 179.47 160.52 2 PLICEION 389 18.30 19.53 14.91 27.77 5.31 8.25 1.50 14.05 18.05 18.05 18.0 18.0 18.0 18.0 18.0 18.0 18.0 18.0	5	PLACE1011353	13.06		9.31							**	- 1	-	- 1
PLACEIDISS S. 4.7 A.0.06 43.45 30.4 28.76 22.51 35.29 36.44 44.51   -   -					7.09								- 1	-	
PLACEIDIST, S. 03 11.85 15.3 3.77 4.12 3.15 3.67 7.88 4.18 \$ -   PLACEIDIST, 7.04 5.27 10.93 2.06 3.1 2.3 1.1 2.3 1.72 3.77 5.21 \$ -   PLACEIDIST, 7.04 5.27 10.93 2.06 3.1 2.3 1.1 2.3 1.72 3.77 5.21 \$ -   PLACEIDIST, 7.04 5.27 10.93 2.06 3.1 2.3 1.1 2.3 1.7 3.77 160.52   PLACEIDIST, 7.04 5.27 10.93 2.06 3.1 2.3 5.0 1.4 0.5 16.53 13.24 \$ -   PLACEIDIST, 7.05 1.3 5.1 4.9 17.7 5.3 1.8 2.3 5.0 1.4 0.5 16.53 13.24 \$ -   PLACEIDIST, 7.05 1.3 5.2 5.14 17.24 17.93 13.0 1 14.05 16.53 13.24 \$ -   PLACEIDIST, 7.05 1.3 6.3 1.3 1.4 4.0 1.2 5.1 1.3 4.3 9.2 4.88 4.7 7.06 4.9 15.2 1.3 1.4 29.6 1 *   PLACEIDIST, 7.05 1.3 6.3 1.3 1.4 4.0 1.2 5.1 1.3 1.5 8.9 3.2 4.8 2.4 8.0 4.0 1.8 3 4.2 2.5 1.3 1.2 24.1 29.6 1 *   PLACEIDIST, 7.05 1.3 1.4 5.6 1.3 1.4 4.0 1.2 5.1 1.1 5.8 9.2 2.3 4.5 2.4 0.0 1.8 3 4.2 2.5   PLACEIDIST, 7.05 1.3 1.1 5.8 1.5 1.1 5.8 9.2 2.3 4.5 2.4 0.0 1.8 3 4.2 2.5   PLACEIDIST, 7.05 1.3 1.1 5.8 1.3 1.4 1.2 1.3 1.4 1.2 1.3 1.4 1.2 1.3 1.4 1.2 1.3 1.4 1.3 1.4 1.3 1.4 1.4 1.4 1.4 1.4 1.4 1.4 1.4 1.4 1.4		PLACE1011364	36.2	48	46.69	18.82	32. 17	16.48	29.76	23.08			*	-	-
PLACEIDISTS 7, 04 6.27 10.33 2.06 3.1 2.3 1.12 3.17 5.21 1 -		PLACE1011365		40.06	43.45	30.4	25.78	22. 53	35, 29				- 1		- [
PLACEID1138   164, 23   196, 07   182, 53   143, 55   164, 65   123, 22   185, 52   179, 47   160, 52   PLACEID1149   135   14, 31   177, 75, 31   182, 35   101, 140, 55   16, 51   12, 24   PLACEID1149   45, 63   174, 47, 01   28, 72   27, 55   28, 25   28, 24, 28   47, 7, 06   48, 28   52   48, 28		PLACE1011371	8.03	11.85	15.3	3.37							- 1		- 1
PLACEID11386   64, 63   160, 07   182, 53   14, 65   124	10											*	- 1	-	- 1
PLACEID11469 24.55 18.26 25.14   17.24   17.92   13.01   14.05   16.53   13.24   2   3   -   PLACEID1471   1.96   8.15   11.8   4.39   4.25   4.88   4.25   4.88   4.89   5   2   -   PLACEID1471   1.96   8.15   11.8   4.39   4.25   4.88   4.25   4.88   4.88   5   2   -   PLACEID1469   27.4   5.05   13.15   5.89   27.15   27.55   25.25   33.12   24.14   29.66   7   -   PLACEID1460   22.91   21.43   16.68   17.18   6.55   6.34   4.22   2.79   3.13   5   5   5   PLACEID1465   2.99   27.18   27.18   5.55   5.3   3.31   2.44   29.66   7   -   PLACEID1465   2.59   27.18   27.18   27.55   25.5   3.31   2.44   29.66   7   -   PLACEID1477   8.38   4.15   8.42   3.82   3.5   4.74   4.12   4.56   4.31   2.75   3.31   PLACEID1477   8.38   4.15   8.42   3.82   3.51   2.48   5.88   2.55   3.31   2.48   2.55   3.31   2.48   2.55   3.31   2.48   2.55   3.31   2.48   2.55   3.31   2.48   2.55   3.31   2.48   2.55   3.31   2.48   2.55   3.31   2.48   2.55   2	10												- 1		
PLACEID11407 11.96 8.15 11.8 4.39 4.52 4.88 4.7 7.06 4.99 12 2   PLACEID11407 11.96 8.15 11.8 4.39 4.52 4.88 4.7 7.06 4.99 12 2   PLACEID11407 12.5 6. 31.74 4.01 28.72 27.55 28.52 31.12 24.14 22.96 6 2 + -   PLACEID11403 12.4 6.05 12.11 5.89 9.23 4.52 4.00 4.22 2.79 3.13 4.28   PLACEID11465 22.91 21.43 16.86 17.18 6.85 6.34 4.12 2.79 3.13 4.28   PLACEID11465 18.59 10.4 8.63 17.18 6.85 6.34 4.12 2.75 3.13 4.28   PLACEID11471 12.35 11.03 8.39 5.76 5.63 4.74 1.72 3.62 2.79 3.42 2.9 18 2.4   PLACEID11471 12.35 11.03 8.39 5.76 5.63 4.74 4.12 4.06 4.30 4.8 1												**	- 1	-	ı
PLACEIDI149  45.6   31.74   47.01   28.72   27.55   28.25   31.12   24.14   29.66   1   2   2   2   2   2   11   5.89   22.3   4.55   4.4   4.22   2.79   3.13     ##   -															- [
PLACEIDI1433   12.4   6.05   13.11   5.89   9.23   4.52   4.03   6.18   4.28   7.18													*		-
PLACEID11440   22.91   21.43   16.85   17.18   6.45   5.14   4.12   27.79   31.13   ## -   PLACEID11467   21.99   27.18   27.18   27.18   5.85   8.31   4.18   2.55   3.16   3.27   3.13   ## -   PLACEID11467   21.99   27.18   27.18   3.15   5.85   8.31   3.18   2.55   3.76   2.55											Z9. 66	*	- 1	-	- 1
PLACEID11452 21.99 27.18 21.97 74.77 5.58 8.31 1.98 2.55 3.76 5.82 4.7 - PLACEID11472 12.35 11.03 8.39 5.76 5.83 4.74 4.12 4.06 4.53 8 8 PLACEID11472 12.35 11.03 8.39 5.76 5.83 4.74 4.12 4.06 4.53 8 8 PLACEID11472 12.35 11.03 8.39 5.76 5.83 4.74 4.12 4.06 4.53 8 8 PLACEID11472 12.35 11.03 8.39 5.76 5.83 4.74 4.12 4.06 4.53 8 8 PLACEID11472 14.78 12.74 5.15 1.77 8.47 3.82 3.10 12.49 5.86 5.38 7.03 8 PLACEID11478 14.78 12.74 5.15 1.77 8.48 12.81 14.29 19.74 4.78 12.22 2.15 1.79 12.	15												1		- 1
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PLACEIDI1472   2.35   11.03   8.39   5.76   5.62   4.74   4.12   4.06   4.53   * **   PLACEIDI1473   47.8   127.45   151.17   88.47   3.82   3.1   2.49   5.68   5.36   7.05   **   PLACEIDI1478   47.8   127.45   151.17   88.41   123.14   82.91   97.94   88.22   67.65   **   PLACEIDI1478   11.85   1.27   11.87   6.42   7.11   5.44   7.49   6.78   2.62   7.10   **   PLACEIDI1503   8.16   7.18   8.72   3.16   3.42   2.13   5.10   9.05   8.10														-	
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## PUACEIOIISS 21.22   18.3   17.27   18.5   16.2   29.2   27.07   30.14   \$2												**		-	
## PULCEIOIISE   56.54   29.6   21.88   17.54   11.85   16.02   29.9   27.07   30.09   2   -   PULCEIOIISE   29.7   3704, 8   30.31   195.59   285.57   285.01   30.07   3775, 5   2747.5   5   -   PULCEIOIIST2   14.56   19.19   17.05   4.82   11.54   6.52   8.71   6.56   7   8.56   8   8   -   PULCEIOIIST2   27.27   28.5   23.64   17.67   7.67   13.02   10.47   6.81   13.88   8   8   -   PULCEIOIIST2   20.54   2															
*** PLACE 1011176   469.7   3704.8   3103.3   1955.8   2495.7   2590.1   3091.7   3779.5   2474.5   8   -   ***PLACE 1011877   27.27   22.5   23.64   11.76   7.61   31.02   10.47   6.83   11.83   ***   -   **PLACE 1011878   27.27   22.5   23.64   11.76   7.61   31.02   10.47   6.83   11.83   ***   -   **PLACE 1011878   27.27   22.5   23.64   17.6   7.61   31.02   10.47   6.83   11.83   ***   -   **PLACE 1011789   240.54   255.47   223.18   186.78   250.35   189.86   147.11   130.97   144.28   **PLACE 10117129   13.64   15.82   1953.8   165.8   9100.2   124.6   880.28   152.8   **PLACE 1011729   14.63   11.44   8.67   4.74   6.3   2.04   9.22   9.94   8.53   *   **PLACE 1011729   13.65   10.92   5.48   6.55   5.70   02.01   24.66   10.55   15.85   5.87   **PLACE 1011741   19.11   16.45   8.67   4.68   8.95   8.58   13.7   33.5   14.66   10.55   15.85   5.6   *   **PLACE 1011757   15.5   13.64   8.67   8.67   8.67   8.37   7.2   13.7   32.7   10.55   6.56   *   **PLACE 1011757   12.55   13.61   8.61   93.67   33.7   32.7   13.5   35.65   8.5   **PLACE 1011776   14.17   14.57   14.58   8.09   8.23   7.54   9.58   8.34   17.29   4   **PLACE 1011778   24.3   3.67   23.05   14.78   12.6   12.7   23.5   14.77   23.5   14.6   10.55   15.6   5   **PLACE 1011778   34.3   3.67   23.05   14.78   12.6   12.7   23.5   14.77   23.9   23.6   14.7   23.9   23.6   23.7   23.5   14.7   23.9   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5   23.6   23.7   23.5													*	-	-1
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PUCEIDITISTS 27.22 28.5 23.64   1.76 7.61 13.02   10.47 6.35 11.88   ** =   PUCEIDITISTS 27.22 28.5 23.64   1.76 7.61 13.02   10.47 6.35 11.88   ** =   PUCEIDITISTS 20.55 25.47 223.18   186.79 250.35   189.86   147.11   130.87   144.28   ** = -   PUCEIDITITIST 21.51 24.5   180.28   1893.8   185.5   100.02   142.6 860.26   126.5   PUCEIDITITIST 1.52 11.6   10.22   5.48 6.5 1.57   9.02   5.48 6.5 1.57   9.6 6.5 1.57   PUCEIDITITIST 1.51 13.6   16.45   8.77   6.48 6.39 4.13 6.03 4.25 6.00   ** = -   PUCEIDITITIST 1.55 11.86   18.61   93.77   137.7   33.5   146.6 10.5 5   16.55   ** -   PUCEIDITITIST 1.55 11.86   18.48   96.85   33.7 57.2   11.37   92.71   10.5 5   * = -   PUCEIDITITIST 1.55 11.86   14.89   66.55   33.7 57.2   11.37   92.71   10.5 5   * = -   PUCEIDITITIST 25.5 11.86   14.98   66.55   33.7 57.2   11.37   92.71   10.5 5   * = -   PUCEIDITITIST 25.8   18.6 1.9 18.71   18.16   17.09   15.47   71.92   2.65   2.65   PUCEIDITITIST 26.85   17.85   14.49   8.65   5.27   6.8   13.7   72.7   72.7   72.6   5.6   * = -   PUCEIDITITIST 26.86   17.85   14.49   8.65   3.7   72.0   72.0   72.0   72.0   72.0   72.0   PUCEIDITITIST 26.85   13.5   13.5   23.5   23.7   23.5   13.5   23.															- 1
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45 PLACEIDITIFS 2115. I 1224. 5 1828. 8 1935. 8 1985. 8 1985. 9 100.2 2 1426. 8 860. 28 1289. 9 PLACEIDITIFS 2115. 1 1224. 5 10.92 6.87 4.74 6.3 2.04 9.22 9.39 4.55 5 = - PLACEIDITIFS 17. 92 12.45 10.92 6.88 6.85 1.57 9.59 6.92 10.12 4 - PLACEIDITIFS 19.11 16.45 8.47 6.48 6.85 1.57 9.59 6.92 10.12 4 - PLACEIDITIFS 19.51 6.48 6.49 6.85 1.57 9.59 6.92 10.12 4 - PLACEIDITIFS 19.57 9.64 1.51 1.3 22.6 27.43 20.99 15. 9 4.76 6.86 4 4.8 4 - PLACEIDITIFS 19.57 6.43 15.13 22.6 27.43 20.99 15. 9 4.76 6.86 4 4.8 4 - PLACEIDITIFS 11.2 55 18.41 84.99 6.65 85.7 75.4 20.95 10.95 27. 109.2 4 - PLACEIDITIFS 11.4 71 4.57 14.58 8.99 8.7 138.7 13.7 13.7 12.7 109.2 4 - PLACEIDITIFS 28.88 17.85 14.48 8.49 16.55 27. 75.6 12.3 20.45 4 1.99 1.99 1.99 1.99 1.99 1.99 1.99 1.												i			
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PLACEIDIT41   19.11   16.45   8.77   5.48   6.39   4.12   6.03   4.52   6.01   2   3   -   PLACEIDIT41   19.11   16.45   18.11   19.17   13.77   39.51   46.78   16.95   16.95   56   2   4   PLACEIDIT57   55.97   64.31   51.13   22.6   27.43   20.99   59.9   47.96   55.48   4   -   PLACEIDIT57   14.17   14.57   14.58   18.95   65.85   37.57.2   11.37   82.71   10.92   2   -   PLACEIDIT78   14.17   14.57   14.58   8.09   8.23   7.54   9.58   8.14   17.29   4   -   PLACEIDIT78   26.85   17.35   14.95   8.49   7.89   4.41   13.15   9.37   14.55   2   -   PLACEIDIT89   26.85   17.85   14.49   8.41   7.89   4.41   13.15   9.37   14.55   5   -   PLACEIDIT89   5.46   2.52   3.15   2.33   2.07   0.8   3.19   3.06   3.38   PLACEIDIT29   26.52   55.93   50.79   29.92   21.65   12.01   21.62   24.55   19.64   8   -   PLACEIDIT29   27.71   53.31   45.03   28.52   44.77   24.15   24.4   44.77   47   8   -							6.00	2.57						-	- 1
PLACEIDITI49 188. 63 245. 38 185. 1 98. 37 138. 73 83. 5 146. 64 106. 55 169. 56 8 - PLACEIDIT75 15. 87 64. 31 51. 13 22.6 27.4 20. 99 59. 9 47. 96 65. 46 ** - PLACEIDIT76 112. 55 118. 61 84. 99 66. 65 83. 7 57. 21 113. 37 82. 71 109. 2 * - PLACEIDIT76 14. 17 14. 57 14. 58 8. 99 8. 23 7. 54 9. 55 8. 47 17. 29 ** - PLACEIDIT78 34. 3 38. 72 30. 51 47. 82 11. 16 17. 99 15. 47 21. 92 20. 45 * - PLACEIDIT79 26. 86 17. 86 14. 49 8. 41 7. 89 4. 41 13. 16 9. 37 14. 56 * - PLACEIDIT98 26. 46 25. 23. 135 2. 32. 70 8. 41 13. 16 9. 37 14. 56 * - PLACEIDIT98 26. 52 55. 33 50. 79 29. 92 21. 66 12. 01 21. 62 24. 56 19. 64 * 9LACEIDIT89 27. 13 3. 14 13. 18 9. 37 14. 56 * PLACEIDIT89 27. 13 13 14 15 15 15 15 15 15 15 15 15 15 15 15 15														-	- 1
PLACEIDITI75 55.97 84.31 51.13 22.6 27.43 20.99 58.9 47.96 65.46 42 - PLACEIDIT76 11.25 11.86 18.49 65.65 38.7 57.21 11.37 82.71 10.92 4 - PLACEIDIT76 14.17 14.57 14.58 18.99 65.25 38.7 57.21 11.37 82.71 10.92 4 - PLACEIDIT76 14.17 14.57 14.58 19.09 8.21 7.54 9.58 8.34 17.29 47 - PLACEIDIT76 25.86 17.36 14.99 8.41 75.99 4.41 13.16 9.37 14.56 2 - PLACEIDIT86 5.46 2.52 3.15 12.86 17.89 4.41 13.16 9.37 14.56 2 - PLACEIDIT86 5.46 2.52 3.15 2.35 2.70 0.8 31.9 3.0 5.0 3.38 19.00 13.88 19.00 10.12 12.00 12													•		- 1
99 PULCEIDITIES 12. 55 118. 61 84. 93 86. 85 83.7 87.21 113. 37 92.71 109.2 2 - PLACEIDITIES 14.17 14.57 14.58 9.09 8.21 7.54 9.58 8.34 17.29 4# - PLACEIDITIES 34.3 38. 87 23.05 14.78 12.16 17.09 15.84 21.92 20.45 * - PLACEIDITIES 26.86 17.86 14.49 8.41 7.89 4.41 3.16 9.37 14.66 * - PLACEIDITIES 6.46 2.52 3.15 2.33 2.07 0.8 3.19 3.06 3.38 * - PLACEIDITIES 6.45 2.55, 33 50.79 29.92 21.66 12.01 21.62 24.55 19.64 * - PLACEIDITIES 4.67 17.15 3.11 45.03 28.52 44.7 24.18 6.14 24.87 87 47 4* - PLACEIDITIES 4.71 15.31 45.03 28.52 44.7 24.18 6.14 24.87 87 47 4*															- 1
PLACEIDIT78 14.17 14.57 14.58 9.09 8.22 7.54 9.58 8.34 17.29 98 - PLACEIDIT78 14.2 15.4 5.2 1.55 14.58 19.09 8.22 7.54 9.58 8.34 17.29 98 - PLACEIDIT89 28.68 17.68 14.49 8.41 7.89 4.41 13.16 9.37 14.56 8 - PLACEIDIT89 5.46 2.52 3.15 2.3 2.07 0.8 3.19 3.06 3.38  PLACEIDIT89 8.62 5.53 50.79 28.92 21.66 12.01 21.62 24.56 19.64 8 - PLACEIDIT89 4.11 53.11 45.03 28.52 14.67 24.01 21.62 24.56 19.64 8 - PLACEIDIT89 4.11 53.11 45.03 28.52 14.67 24.17 24.18 6.12 4.48 74 48	50														- 1
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PLACEI011810 5.46 2.52 3.35 2.53 2.07 0.8 3.19 3.06 3.38 PLACEI011824 8.6 2 55.93 50.79 25.92 21.66 12.01 21.62 24.55 19.64 \$\$ \$\$														۱-	- 1
55 PLACEI011824 86.62 55.93 50.79 29.92 21.66 12.01 21.62 24.56 19.64 \$ \$ PLACEI011825 47.11 53.31 45.03 28.52 34.47 24.16 61.42 44.87 47 ** -												1		1	- 1
PLACE1011825 47.11 53.31 45.03 28.52 34.47 24.16 61.42 44.87 47 ** -				55.93										-	- 1
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	PLACE1011836	18. 92	14, 13	16.67	5.56	9.72	5. 44	13.34	9. 9	14.01		- 1		- 1	
	PLACE1011847	18, 99	16.08	12,62	9, 39	11.06	5, 84	18.15	10.26	13.1		- 1	-	- 1	
	PLACE1011855	44, 57	41, 99	25. 3	17.11	33, 33	14, 76	55.6	42.68	40.57		1		- 1	
5	PLACE1011858	17.26	16.38	17, 11	5.33	7.08	6, 53	6. 22	8.74	5.1	**	**	-	-1	
	PLACE 101 1874	23, 12	12,72	25.71	16.84	19.22	7.85	10, 17	8.71	12.51		- 1		- 1	
	PLACE 1011875	10, 58	9, 98	11, 17	5.18	10.63	6.87	7.18	5.04	4, 93		**		-1	
	PLACE 1011877	4, 39	6.14	4. 95	3.7	5, 52	3,78	2.59	4.05	4, 24		- 1		- 1	
	PLACE 1011891	21.71	17.57	18.99	7, 39	14, 47	8,67	8, 21	9, 15	7.86		**	-	-1	
	PLACE 1011896	15, 35	16.64	24.3	10,41	10, 14	12, 14	5, 01	8. 13	7. 98		*		- 1	
10	PLACE 1011920	16, 12	14, 42	15. 9	7.68	8, 85	7.76	6, 46	2.46	3, 24	**	**	-	-1	
	PLACE1011922	9, 14	7. 55	8.03	4.49	8. 39	4, 08	7.97	6.15	5. 33				- 1	
	PLACE1011923		139, 43	146.9		117.42	86. 95	136. 93		124, 61		- 1	_	- 1	
	PLACE 1011937	11. 45	9, 18	10.8	8.58	6.75	5. 1	5, 71	6, 29	5.46		**	-	-1	
	PLACE 1011939	20. 33	21.55	26.93	12.79	15. 19	13, 44	14, 58	11, 31	12.67			-	- 1	
	PLACE 1011940	46.38	48.76	46.73	35. 11	34. 53	31.67	43, 31	41.44	39. 86	**	**	_	-	
15		58, 59	70.73	80.18	30.15	38.09	28.84	47. 96	47.68	42.57	**	7	-	- 1	
	PLACE1011962 PLACE1011964	53.78	49.36	46, 95	41, 43	47, 33	36.34	22. 23	20.62	20.74	**			-1	
		50, 21	52.86	64.71	38.6	31, 39	22. 36	22.46	20. 19	18, 12			_	-	
	PLACE1011978						6, 53	4, 68	9, 99	8. 31	•	"	-	- 1	
	PLACE1011980	13. 57 46. 23	18.54	10. 96 45. 44	9.65	15, 28 15, 61	9, 98	14.66	11.01	11.18		**		- 1	
	PLACE1011981		28.89		16, 98	68.6	45, 01	97.45	90.55	115.07	**	**	-	-	
20	PLACE1011982	116.69	11.27	123. 56 12. 33	57. 42 5	7, 29	7. 22	6, 27	8.29	4. 87	7		Ξ.	•	
	PLACE1011995	8.31 36.36	36.13	39. 16		26, 03	23. 3	20.6	21.06	24.79	**	**	-	-	
	PLACE1012023				21. 46		50.42		33.87	25. 71	**	**	_	-1	
	PLACE 1 01 2026	72. 96	80.46	66, 75	52.71	40. 29		21.72	3.11	3. 97	**	"	-	-	
	PLACE1012031	6.69	7. 34	9.86	2.73	6.11	3.89	3.41	32, 21	30.64		٠,		-	
	PLACE2000003		222.75	62.64		133.88	40.79	39. 01	8. 65	14.3				-	
25	PLACE2000005	20.72	25. 78	18. 16	9.79	22. 43	11.52	5.79				*		-	
	PLACE2000006	74. 92	51.78	61.3	27,4	24. 75	21.12	50.02	43.94	72.77	**		-	-	
	PLACE2000007	11.31	11.29	12.53	6.68	7. 02	5.2	5.63	10.02		**	•	-	-	
	PLACE2000011	20, 07	19.36	23.84	14. 92	4. 58	15.77	9, 39	11.52	19. 43					
	PLACE2000014	10.47	6.72	12. 3	4.76	6.02	3.48	3.63	6.9 26.64	4.73	*		-	-	
	PLACE2000015	63.77	83.22	43.62	34.06	43.14	43. 39	19.29		36.8		•		-	
30	PLACE2000017	64.44	66.07	78.07	40.68	48.09	29, 95	59.7	61.53	44. 91	*		-	-	
	PLACE2009021	131				142.66	101.82	89.75	68.71	86.35		**		-	
	PLACE2000022	6. 52	6.75	7.52	3. 85	5, 14	3.55	4. 53	1. 84	5. 72	**		-	_	
	PLACE2000030				182.81		156.31	125. 23	96.07	135.65		**	-	~	ı
	PLACE2000032	8, 03	6.4	8.8	4, 38	5. 33	2.81	4. 32	15.18	4. 01	*		-	-	
	PLACE2000033	19.24		20.22	9.43	14.03	10.83	8.93	12.91	10.87	*	**	-	- 1	
35	PLACE2000034			146.24		195.88			112.34			. 1			
	PLACE2000039	8. 9	10.84	13.51	4. 94	3.89	4. 8	3.72	4. 38	6. 52	**	*	-	-	l
	PLACE2000043	13.64	20.44	15.11	6. 28	13. 12	6,04	6. 05	7.64	6. 22		*		-	
	PLACE2000044	7.28	9. 48	9.44	3, 48	5.71	4.79	4, 48	4. 25	4, 59		**	~	-	ı
	PLACE2000047	17, 41	19.39	12.83	6.83	12.83	7.05	7.81	8, 54	5.78		*		-	ı
40	PLACE2000050		17.87	22.05	7. 34	8. 98	7.31	7.51	8. 31	7. 94	**	**	-	-	
40	PLACE2000061	53.75		78. 23	34. 92	41.03	37.86	22.68	14.04	20.77	*	**	-	-	ı
	PLACE2000062	44. 19	44.8	62.08	41.54	45. 17	33.58	38.88	38.06	40. 24				-	
	PLACE2000072			181.98					113.44	105. 53		**			
	PLACE2000073	8.16		9. 27	3.77	4.89	3. €2	4.56	4. 79	3.59	**	**	-	-	
	PLACE2000097		128.07			131.74		90.14	87.25	86.79		**		-	ı
45	PLACE2000100			15.99	6.02	8.77	5.56	4.8	3. 51	6.24	**	**	-	-	ı
45	PLACE2000103					436. 52			336.03				1		ı
	PLACE2000106			23.14	14.46	6.17	13.03	9.13	10.5	5. 27	*	**	-	-	ı
	PLACE2000111		157.38	125. 3		105. 91	83.66	90.52	44, 23	69.19		**	-	-	ı
	PLACE2000115			16.32	19.6	21.67	16.26	13.74	11.32	11.63			1		
	PLACE2000118			143. 27	124. 01		92.15	153.7	62.85	67.07					
50	PLACE2000124			5. 57	2. 12	1.78	1.55	2.01	1. 56	1.09	**	**	١-	-	
30	PLACE2000132			19.11	15.64	11. 16	12.49	9.7	6. 52	8. 48	١.	**	-	-	ı
	PLACE2000136			26. 33	13.95	42.68	22.54		17.8	18. 28	۱		1	-	ı
	PLACE2000137			6.91	1.61	2. 95	2.09	1.24	1.8	2.36	**	**	-	-	ı
	PLACE2000140			12.2		9. 45	9. 49	11.48	11.25	10.8	**		-		ı
	PLACE2000147					5.85	4.51		7.6	9. 27	*		-		ı
55	PLACE2000153			13.85		9.57	6.43	4.84	5. 27	2. 27	*	**	۱-	-	ı
~	PLACE2000164			9. 44			4. 12		3. 97	2. 13	**	**	-	-	1
	PLACE2000170	38.84	46.93	33	28	28. 56	22. 27	30.03	23.36	30.82			-		ı

						able 4	94								
	PLACE2000172	9, 55	12.02	10.86	4, 88	5, 87	3. 32	4.01	4.8	6.2	**	**	-	- 1	
	PLACE2000173	16.63	20,86	19,04	9.66	11.13	9.7	6.74	6.23	3, 14	**	**	-	- 1	
	PLACE2000174	6.73	7.76	6.4	3.37	4. 73	2.4	3, 08	3.26	1.62		**	_	- 1	
5	PLACE2000176	47.07	39. 92	36.01	26. 8	32.78	24. 97	24.68	26.79	25, 41	i	**	-	-1	
3								2.47	4. 64	3, 94	ï	: 1	_	-1	
	PLACE2000187	11.49	9.44	7. 22	4, 28	4. 58	4.45		4. 64			:	-		
	PLACE2000216	15.24	13.28	16.1	7.83	10. 53	7.67	11.12	11. 23	10.67	**		-	~	
	PLACE2000219	43. 32	28. 21	37. 3	18.17	6.67	13. 94	7.68	7,79	6.4		**	-	-	
	PLACE2000221	9.58	7.64	7.46	2.22	3, 99	1.24	4.08	2.43	3.67	**	**	~	~	
	PLACE2000223	16.67	12.34	12.95	4. 95	6.75	4. 32	4.43	2.01	2.87	**	**	-	-1	
10	PLACE2000231	8, 48	8, 65	7.01	3.78	5, 92	4. 27	4, 19	2, 32	3. 59	*	**	-	- 1	
	PLACE2000235	7.02	5.03	3.66	1.54	5. 33	1.9	4.43	3, 57	3. 78				- 1	
	PLACE2000246	21.31	17. 8	13. 21	7.9	13.4	6. 47	11.52	9.6	10.91		.		- 1	
				30, 68	20.28	22. 27	18. 14	15, 54	15.85	15.69			_	-	
	PLACE2000264	41.11	43.3					3. 84	3, 12	4, 31	÷		_	- 1	
	PLACE2000274	6.8	7.42	7. 36	4. 35	3.04	1,89		15.3	19.45	••		_	-1	
15	PLACE2000287	32.11	30.76	24. 99	20.39	25. 61	21.53	17.96				** [		~ [	
	PLACE2000296	42. 94	34.88	35.63	23. 36	22.17	20.37	32.02	14, 36	29. 35	**	)	-	- 1	
	PLACE2000302	252.74	304. 77	237.45	265.35				112.58	108.48		**		- 1	
	PLACE2000305	21.47	23.58	17.69	16, 14	17, 23	16.65	22.14	18. 21	22.01		i		- 1	
	PLACE2000317	37.2	38, 12	23.24	20.01	28, 38	14, 37	31.05	33, 16	31, 64		- 1			
	PLACE2000324	18.44	17.84	10.35	6. 13	8.75	4, 98	3.24	8.74	3.2		* 1	-	- [	
	PLACE2000334	21.38	17.6	11.43	10, 36	9.7	9. 37	7. 54	6, 15	9. 1		* I		-1	
20	PLACE2000335	71.59	66. 59	54.82	51.15	35, 99	40, 86	33.77	36.69	29.67		##	_	- 1	
		20.62	28.95	19	17. 88	23. 88	12.85	16. 23	14.73	16. 82		1		- 1	
	PLACE2000340										**			-1	
	PLACE2000341	14.8	11.92	10.45	5.66	5. 34	5. 33	5. 38	5. 46	7. 66		٠,	_	- 1	
	PLACE2000342	23.53	21.95	21.97	16.28	11.86	11.51	22.81	14.69	23. 12	**		-	- 1	
	PLACE2000347	9. 57	9. 26	10.05	4, 16	2. 94	3, 3	5.78	3.54	5. 48	**	**	-	- 1	
25	PLACE2000357	112.17	70.59	69.73		100.62	78.25	23	68.79	48, 54		i		- 1	
	PLACE2000358	57. 37	32.59	34.74	25.22	20.69	15.46	44.93	47.18	42.01				- 1	
	PLACE2000359	66.39	48.68	57.2	40.91	54, 54	32.45	1.96	27, 1	55				- [	
	PLACE2000366	24. 09	23.36	19.56	12.65	12.71	9. 05	11.75	9, 72	10.83	**	**	-	-1	
	PLACE2000371	12, 17	9. 37	8, 13	5, 98	7.76	4.49	7.64	10.97	6. 2		1		- 1	
	PLACE2000373	8.3	7.58	8, 83	4, 52	6. 25	4, 81	3.89	1, 17	2.44	**	**	-	-1	
30	PLACE2000374		21.57	18.21	16.3	16.71	14. 02	11.71	11,44	14. 19		**	-	-1	
30				125. 02	99.68	141.5	89. 14	84.64	65, 75	71.64	•	**		-1	
	PLACE2000379		110.55						17.99	10.26	**	-		-	
	PLACE2000386	27.43	27.19	20.99	10.83	12.4	11.22	15.3		93.57	**		_	-	
	PLACE2000388			203.4					109.84						
	PLACE2000392								115.9	95.8		**	-	-	
	PLACE2000394	10.9	10.42	13.61	8.59	15. 25	7.68	5.68	9.12	8.51		*		-	
35	PLACE2000398	16.83	19.82	17.04	9.21	12, 98	10.99	7.5	15. 14	8.57	**	*	-	- 1	
	PLACE2000399	30. €	27.17	17.03	25.83	27.41	28.06	9.08	19.74	20.17				- 1	
	PLACE2000402	5.25	8.04	8. 31	4.2	9.81	3.39	4.84	3.89	6.41				- 1	
	PLACE2000404	24.79	49, 52	29, 14	49.74	21, 66	25, 22	1€.23	19, 08	8, 51				- 1	
	PLACE2000411	5.67	8, 96	6.76	7.21	11,84	6, 13	5.19	4, 54	7.08				- 1	
	PLACE2000418	10.3		9, 67	6.1	3.09	4. 67	4, 68	5, 3	4.11	**	**	-	- 1	
40	PLACE2000419	96.71	94, 48		58.48	57. 97	27.06	44.37	46.85	45, 47			_	-1	
	PLACE2000425	9, 83		13, 49	7.49	11. 26	6.24	5, 84	7.49	5. 62		*		-1	
	PLACE2000427	15.21	22.92	21.67	15.7	21.56	15. 92	5.61	6.9	4.76		**		- 1	
	PLACE2000427	25.58		26, 12	13.97	18.4	14. 15	21.62	27.64	18,08	**	**		1	
											1			-1	
	PLACE2000435	16.36		21. 19	9.69	13.76	7.6	8. 43	10.79	12.73	١.	٠,	-	-1	
45	PLACE2000438				24.14	34. 38	15, 98	38. 19	37.23	25. 32			ı	- 1	
45	PLACE2000450		41.59		16.35	37, 33	18.21	23.47	16.65	30.25			l	- 1	
	PLACE2000455	51.05			30.31	36, 57	21.71	39.12	34. 02	35. 42		**	-	-1	
	PLACE2000458	13.07	11.58	13.21	6.82	7.75	7.03	6.6	8.78	5.76	**	**	-	-1	
	PLACE2000464	9.95	10.55	11.94	7.59	6. 37	5, 51	7, 66	8.47	7.16	**		-	-1	
	PLACE2000465	182.2			166.02	148, 04	156.91	139.29	234.97	151.61			-	- 1	
	PLACE2000473	23.66		24. 43	18.65	18, 15	16.44	24.18	30.07	29. 1	**		-	- 1	
50	PLACE2000477				9.69	12.47	6.23	10.6	9. 09	9, 9	1		ł	-1	
						53.35	39.93	39.48	28, 43	32.45	i	÷	1	-	
	PLACE3000004									20. 29	1	**	1	-1	
	PLACE3000009					35.05	20. 31	13.54	15.44		١		1		
	PLACE3000020					15.64	8, 65	14. 29	14.44	13.63	**	**	1 -	-1	
	PLACE3000029					9. 27	7.3	4. 68	16.59	5. 52	**		-	- 1	
55	PLACE3000038	6. 52	4. 32			3.97	4. 12		6.95	6.18			ı	- 1	
~	PLACE 30000 52			10.48	3.53	4, 29	4, 15	2.75	5.07	2.29	1		1	- 1	
	PLACE3000059					5.08	2.55		4. 28	4. 31		*	-	- 1	

							Table 4	195							
		000067	49.68	45.52	57. 37	38.78	41, 13	26.16	39.8	46.08	41.69	l		l	- 1
		1000069	22.6	20.15	23	9, 32	14.75	10.81	1.9	9.3	14.16	**	*	-	-
		000070	28. 36	29. 24	22.17	14.93	25. 34	18.73	8. 92	18. 55	16.21		*		-1
5		1000103	7.35	7.9	9. 44	5. 25	4.6	6.13	2.42	4. 32	5.29	*	*	-	-1
		000119	35. 89	28. 1	30: 76	15.16	14.79	15.09	26.05	30.71	30. 91	**		-	- 1
		1000121	42.11	31.92	37. 82	38, 13	38. 16	27.02	30.65	23. 15	27.55				
		1000124						282.09	411.56	374.59					- 1
		000135		25.87	26.77	15.03	9.64	8. 12	18. 24	11.89	14.6	**	*	-	-
10		000136	7.68	8. 11	13.16	2.72	3. 39	4. 18	4. 52	3.19	1.88	*	*	-	-
		1000142 1000145	24. 15 16. 45	28. 81 12. 13	26. 5 16. 32	11.04 7.23	24. 59	15.08	9. 37	6. 94 9. 12	11.56 6,76		**		-1
		3000147		62.46	138. 9	40.47	16, 52 87, 42	6, 93 32, 89	8. 46 57. 71	63.5	68, 14		*		- [
		000148		59.97	67.42	62.74	65. 96	38. 34	37.63	26.14	36, 79		**		
		000154	10.3	11.14	12.88	7.95	8.64	6. 56	7.31	6.3	7, 57		::		-
		3000155	54.28	50.77	22, 77	23.49	28, 34	24. 7	10.12	11.93	8.34	•	ï	_	-1
15		1000156	9.77	6.45	8.44	2.67	5. 36	4. 07	2, 62	2. 16	2.62		i	_	-
		000157	38.44	50. 52	36, 49	23, 64	29, 36	25.04	21.36	24. 91	20.01		*	_	- 1
		000158	10.22	9. 34	10. 13	4, 45	4. 83	3, 36	2. 93	2. 25	3.02	**	**	-	-
	PLACES	000160	19.92	29.65	15. 93	8.71	11.7	10, 32	5.77	7.29	4.57	*	*	-	-
		1000169	4.65	6.89	7.95	4.88	4.57	2, 21	2.88	2.72	3, 49				-
20	PLACE:	1000181	12.63	9. 87	13.46	6.15	6.69	4. 33	5. 58	7.04	5. 47	**	**	-	-
20		1000194		2636.2	1913.4	1710.4	2606.8	2399	3320.1		2357.1				- 1
		3000197	7.18	4.55	8. 38	2.14	1.71	1.66	2.79	2.24	2.57		*	-	-
		000199	19.46	15.17	17. 17	4.98	8.73	7.24	6. 33	5. 56	6.07	**	**	-	- 1
		000205		16.93	16.84	6. 91	6. 83	4.67	7.28	5. 87	4. 15	**	**	-	-
		3000207	7.97	6,69	7.83	2.48	4. 45	2.85	2.71	2.19	2.08	**	**	-	-
25		3000208 3000213	5. 49 40. 02	5. 5 27. 48	6.02 31.62	1.68 27.29	6. 49 30, 91	· 2. 8 23. 89	2, 64 26, 59	3. 52 30. 58	2.46 23.2		**		-
		1000215	20.76	15.1	17.45	10.19	8, 64	8, 25	15.48	14, 18	17.02	**			- 1
		1000218		10.52	10.44	6.03	3. 81	4. 29	4,65	4, 57	4.68	**	**	-	.
		000220		56.02	42.84	39.7	37	28. 53	36. 15	41.5	44, 54	*	•••	-	- 1
		3000221	12.37	9. 42	8. 54	5.7	3, 63	2,33	5, 09	3, 54	2. 87		**	_	- 1
30		3000225	180.89		131.66	176.14	175. 29	127.3	77.77	64.65	86.2		**		-
-	PLACE	3000226	20.28	21.59	18,74	7,11	11.09	6.3	3.48	2, 23	3.79	**	**	-	- 1
		3000230	17.46	18.98	14.94	6.27	14.11	10.52	4.65	3.25	5. 88		**		-
		3000231	17.8	13.66	8.65	9.08	2.29	4. 95	6.33	9. 35	7.91				- 1
		3000235	9.61	7.55	8. 28	4, 48	5. 65	4.02	5.72	2.93	4.77	**		-	- [
		000242	58. 11	27.4	48. 85	17.82	13.76	17.4	11.61	10.29	11.25	*	*	-	- [
35		3000244	18.63	13.16	20	8.5	7.54	7.64	10.16	10.96	16.61	*		-	
		3000253 3000254	56.99 27.82	47. 97 25. 4	42.85 19.01	32.37	25. 93	24.8	32.98	24.76	32.96	:	*	-	-
		3000254	14.58	20. 8	13.28	15.47 5.71	10. 42 8. 52	14.64 5.71	15. 57 8. 32	11.42 6.43	13.74 8.38	:	:	-	-
		000276	13.09	13.68	8.83	2.78	7.9	3.42	3.8	5. 28	3.1	:	**	-	-
		3000304	37.31	49.14	39. 5	28.72	17.9	18. 26	33, 17	28. 5	40, 23	•	**	_	
40		3000309	12.29	9.64	9.42	6.2	5.6	4. 27	5, 42	4.79	6, 47	**	**	_	-
	PLACES	3000310	51.34	49.06	45, 31	11.5	35, 47	14.78	19.66	9.79	16.36		**	-	-
		000320	24.3	28. 51	23. 33	14.31	20, 51	12.27	15.7	15, 03	15.45	*	**	-	-
		3000322	43.24	35.12	33.62	31.55	31.68	23.75	27.52	20.55	23.56				-
		1000330		27.71	24.49	12.33	15.5	11.35	20.66	17.37	21.33	**		-	-
45		3000331	6.68	8.07	7.67	6.55	5. 13	3,84	2.57	2.99	0.04		**		-
45			393.24	473.08	270.29	359.87		306.41	470,46	662.39					- 1
		1000339		5.98	6. 82	3.79	5. 41	2.48	8.74	10.29	11.53				- 1
		3000341 3000350	12.2 13.71	6.88 12.66	6.99	4.2	5. 65	4. 02	4. 35	5.75 4.37	4.94 7.49				- 1
		3000350	17.23	29, 11	10.21	4. 86 9. 42	8. 13	4.84 8.16	5.77 11.43	7.14	13.58	Ŧ	*	-	-
		3000352	14.74	14.96	16, 82	8.78	15.27 8,26	6.98	10, 37	5.48	6, 79	**	**		-
50		BD00362	57.42	93.8	72.53	\$7.02	59. 59	35. 38	59.68	38. 27	60.6	••	**	_	1
		3000363	9.44	8,61	10.55	4.39	5, 19	4. 3	7.06	4. 68	6, 56	**	*	l _	-
		000365	5. 59	7.45	4. 94	1.48	2.65	3. 2	4.78	2, 37	2.95		•	-	
		3000373	38.45	30. 27	25. 57	4.05	7.88	3, 63	14.16	11. 37	15,74	**		_	- 1
		000374	48.65	44.25	34. 93	16.2	26.01	11. 78	24.23	27. 25	23. 51			-	-
	PLACE	000387	5. 55	3.77	3. 38	1.58	3. 57	0, 43	3.58	2. 95	3. 21				
55		000388		22.21	15. 9	9.86	12.79	6.37	21.94	26.7	24.22	*	-	-	
	PLACE:	000399	15.37	13.07	9.83	5.64	4. 9	3. 93	10.68	11.05	9. 52	**		-	
															-

					1	Table 4	96							
	IPLACE30004001	42, 78	33, 52	30, 29	11,43	26.51	8.471	52,76	33, 47	38, 46	*	1	-	1
		137.51	137.03	95, 06	91.28	111.28	51.38	81.66	72.2	62, 65		*		- 1
	PLACE3000402	8.43	7.72	4, 24	2.44	6, 58	2. 29	8. 31	7.96	5.44		- 1		- 1
5	PLACE3000405	7.33	4.67	7.44	2.73	3.5	5. 1	3.7	5.44	3.66				ı
	PLACE3000406	4. 67	3.39	8. 42	4.46	6.89	4. 16	3.34	4.69	2. 59				- 1
	PLACE3000413	12.33	14.38	16, 34	10.31	15, 31	7.77	6.83	11, 34	9.07		*		-1
	PLACE3000416	84.6	58.17	41.71	94.16	126.08	86, 52	51.61	89.65	63.01				- 1
	PLACE3000425	2, 87	4.19	4.03	2.67	4, 11	2. 12	2.08	0.77	1.54		*		-
	PLACE3000437	6.08	4.62	9.74	3.57	11.5	3	3. 57	3.67	4.77		1		- 1
10	PLACE 3000455	6, 66	7.45	7, 23	5.48	5. 38	4, 19	5.5	5.74	3, 82	*	*	-	-
	PLACE3000475	4. 99	6.71	5.08	4.69	5, 48	5. 91	3.2	5. 35	2.87				- 1
	PLACE3000477	12.61	9.21	10.6	8.81	9.19	10.36	8. 13	7.94	5. 96		*		-1
	PLACE4000003	30	20. 47	39. 94	8. 43	18.43	9. 82	12.11	10.52	8. 23	*	*	-	-
	PLACE4000008	8, 19	8. 19	9. 59	5. 97	7.85	6. 93	4. 18	5. 57	3.47		**		-1
15	PLACE4000009	46.86	36	37. 99	37.22	51.39	35, 97	11.09	18.85	7.53		**		-
	PLACE4000014	18.34	18. 57	19.39	16.62	19. 49	16.59	6.08	13.2	15.87				ı
	PLACE4000029	5.3	5. 07	5. 95	2.91	4.87	2.62	3.69	3.02	2.87		**		-
	PLACE4000034	10.02	8. 01	6. 28	5.12	7. 93	4.55	5. 88	5. 27	5.14				- 1
	PLACE 4000049	21.86	18. 28	18.64	14. 38	19.7	19.05	12.54	13.96	11.06		**		-
	PLACE4000052	47.63	33.03	40.39	17.99	15.08	20. 28	24.01	22.07	24. 29	**	:	Ι.	-
20	PLACE4000062	53.46	46.6	50.02	32.69	40.48	28.08	39, 71	36.99 60.53	45 44. 9		•	-	-1
	PLACE4000063	56.25	58. 52	67.5	55.27 10.66	69, 37 10, 48	73.66	45.78 5.68	5. 4	5.66		**		-
	PLACE4000089	10.35 15.33	13.86 18.02	13.41	10.49	13.84	9. 25 10. 33	17.17	14.6	14.04		**	_	- 1
	PLACE4000093	8. 21	7. 32	11, 21	7.61	6. 2	7, 32	5. 19	5. 2	4, 71	٠.		_	-1
	PLACE4000103	13.85	15.73	12. 84	8. 67	14, 38	6, 43	4.02	8.73	6, 65		**		- 1
25	PLACE4000105	32		23. 23	18.35	21. 38	15, 86	17.78	20.58	18.06		**		- 1
25	PLACE4000128	22.05	18.53	23. 14	8.88	10. 24	9.57	11, 18	17.49	11.76	**		_	-
	PLACE4000129	266.34		188. 25	120.37	95. 41	112, 42	146.64	98.31	136.33	*		-	ı
	PLACE4000131	59. 1	45, 51	65.57	41.68	34. 82	39, 78	18.29	19.7	30.83		**	-	- 1
	PLACE4000147	7.89		11,76	3.51	5. 07	3, 96	2.77	6.32	2. 52			-	- 1
	PLACE4000156	15,48		19. 37	9.1	12. 85	10.65	15.8	15, 48	10.02	*		-	- 1
30	PLACE4000175			18, 08	11.94	9, 67	2.46	6, 38	6.69	6.89		*	1	- 1
	PLACE4000190			32, 85	23.06	35, 22	26.66	12.42	10.73	- 11		**		- 1
	PLACE4000192	35.69	24, 13	23.88	16.77	21.59	17.55	17.75	19.51	19.4			ı	- 1
	PLACE4000206	33.82	29.03	26, 48	11.73	17.97	14.5	17.45	20.16	20.02	**	*	-	-
	PLACE4000211	12.98	13.88	13.86	7.96	9. 94	9, 19	6.64	31.04	6, 54	**			
	PLACE4000214			9. 29	3.45	7.5	4. 16	4.96	7.25	13.71				- 1
35	PLACE4000222			34, 54	24.03	24.44	20. 1	20.58	28.37	25. 45	**	*	-	-
	PLACE4000223			15.03	7.36	10. 28	5. 52	4.16	7.67	5. 85	**	**	-	-
	PLACE4000229			12.52	4.51	6. 31	4.98	4.32	4. 28	3.98	**	**	1:	-
	PLACE4000230			9. 56	4.68	7.06	5. 7	4. 97	9. 82	3.71			1 -	- 1
	PLACE4000233		24. 01	29. 85	11.57	13. 98	13.77	8.53	11.22	9. 26	**	**	1-	-
40	PLACE4000239			9. 42	6.8	7. 34	6.09 8.87	4.49 8.67	4. 54 6. 24	3.54 7.39		**	1	-1
40	PLACE4000247			15. 56 20. 25	9.41 16.54	13. 91 17. 86	12.73	10.47	8.04	11.21	١	**	1	-
	PLACE4000252			12.75	1,39	3.96	3.1	2.21	3. 92	2.7	ı	7		-1
	PLACE4000252			16. 16	12.78	19. 39	8. 44	8. 29	19.39	9.08		•	l	- 1
	PLACE 4000261			22. 9	11.05	14. 2	9. 22	13.61	14. 62	13.29		**	۱-	-1
	PLACE4000264			119.76	135.7		147.27	105, 16	95. 1	89.72				-1
45	PLACE4000265			62.08	23, 03	41.63	24.74	71.71	82.6	52, 18	l .		۱-	- 1
	PLACE4000270			8.9	4.69	5. 92	4.81	3.06	3, 68	3, 44	۱.	**	۱-	-1
	PLACE4000281			18, 29	6,09	10.92	7.61	6.44	6.66	5, 1	**	**	۱-	-
	PLACE4000300			8. 78	6.3	8. 5		3, 68	2.93	2. 96		**	ı	- 1
	PLACE4000320			11.92	6.27	10.09	6.43	4.47	4, 26	2.85		**	۱-	- 1
	PLACE4000323			19.16	9.12	14.06	10.9	6.05	4. 54	4. 03		**	l -	-
50	PLACE4000326			10.53	4.35			4.28	3.68	4.65		**	۱-	-1
	PLACE4000344	27.71	32.93	24. 51	14.27	30.44	19.64	5.14	9.78	8.98		**	ı	-
	PLACE4000347			154. 28	89.79	129.04	79.82	138.06	137.09	135.75	ı		1	- 1
	PLACE4000354	15.47		11, 43	7.01	8. 32			5. 96	4.83		*	ı	-1
	PLACE4000367	8.63		7.82		5. 39			3.95	3. 64	1	*	1	- 1
55	PLACE4000369			9. 31	5.27	4.5			4. 32	3		*	I -	-1
	PLACE4000379								3. 32	2.21		**	1-	-1
	PLACE4000387	25.99	34.88	32.06	17.28	20.68	16.67	6.75	8. 86	3. 97	1 *	**	1 -	- 1

					ı	able 4	91								
	PLACE4000392	8.31	9.76	11.03	3,63	5.95	3.64	5.73	2. 01	2.93	**	*	-	-1	
	PLACE4000399	53. 9	51.54	53.26	20.64	28. 55	18.63	37.94	35.1	44.54	**	**	~	- [	
_	PLACE4000401	4.22	3.76	3.44	3. 96	3.49	2.6	3.5	3.9	1.61		- 1		- 1	
5	PLACE4000403	8.42	5. 82	6.5	4.19	7.67	4.85	3. 96	3. 29	3.53		*		-1	
	PLACE4000411	20. 23	15. 5	15.73	8.23	11.19	9.81	14.98	10.1	11,34	*		-	- 1	
	PLACE4000415	8.15	4.8	4.61	4.06	6.07	4. 23	2.98	3.98	4.5		- 1		ı	
	PLACE4000416	35.43	25.11	24.69	17.72	17.34	16. 23	24.55	17.6	25. 73	*		-	- 1	
	PLACE4000424	43.93	33. 38	31.83	13,69	19, 37	13.98	27.93	23.64	33	**		-	ı	
10	PLACE4000431	10.44	8.78	8.78	5.88	4.85	3.5	4. 69	2. 33 33. 11	4.08	**	**	-	- 1	
	PLACE4000443	50.64	44. 47	41. 43	26, 49	37.08	21.33	51. 22 2. 23	3.95	2.66	•	- 1	~	- 1	
	PLACE4000445	5, 42	4. 17	3.6	3.57	7.43	5. 38 4. 07	3.56	7.58	3.73		- 1		- 1	
	PLACE4000450	9. 85	6.68 132.51	6. 23 128. 81	4.72 98.87	6.64 124.22	95, 79		139.08					- 1	
	PLACE4000455	31.84	25.74	18.83	10.79	23. 42	11. 27	11.29	10.8	17.69				- 1	
	PLACE4000465	13.01	11.54	9. 57	6. 56	6. 92	4, 59	5, 76	4, 65	5.6		**	_	-1	
15		165.28		152. 44	150. 34									- 1	
	PLACE4000472	10.38	6. 99	9. 12	4, 68	4, 63	2, 93	3, 52	2.06	3.49		22	-	- 1	
	PLACE4000489	6.8	7.97	5. 85	3. 99	5.8	4, 43	4, 22	4.11	4, 53				- 1	
	PLACE4000494	70.51	55, 91	43. 58	29, 17	31.84	24.64	72.67	73.79	78,08	*	- 1	-	- 1	
		204.58	180.8	176.1			118.92		165.78	200, 53		1		ı	
20	PLACE4000521	28.26	22, 58	17, 07	7, 12	15.76	9	21.36	20.06	19.61	*		-	- 1	
	PLACE4000522	18.46	20, 38	16.27	10.46	14.51	9.51	16.91	13.88	17.23	*		-	- 1	
	PLACE4000537	9.79	11.19	9.86	4, 99	8.2	4. 43	2.5	3.6	0.68	. *	**	-	-1	
	PLACE4000548	12.35	10.52	9. 27	3.9	4.07	4, 41	4.08	3.04	3.64	**	**	- 1	-1	
	PLACE4000558	56.63	61.59	46, 82	23.74	32.06	21, 71	66.44	39. 24	52.93	**		-	- 1	
	PLACE4000581	9, 34	8. 43	8.7	5. 21	3.71	3. 17	5.23	4.69	4.75	**	**	-	- 1	
25	PLACE4000590	4.28	4. 39	2. 54	2, 29	1.9	1.58	2,69	1.47	1.72	*		-	- 1	
	PLACE4000593	9.43	8. 29	6. 55	6.59	7.86	5. 28	9. 13 8. 58	5. 16 0	4. 1 9. 68		**	١	-	ŀ
	PLACE4000612	50, 54 8, 47	44. 64 6. 12	46.86 6.11	32.73 6.07	32. 12 5. 18	18.75 3.4	1,44	4, 91	1.37	١.		-	-	
	PLACE4000638 PLACE4000650	11.71	6. 12	6. 58	1.78	5. 15	2.69	5.01	3, 89	2, 3	1	•	l		
	PLACE 4000651	10.02	10.01	7.86	4.63	8. 67	3.43	7.21	6.04	5.23	1	*		-	
30	PLACE4000654		37.09	28.74	21.7	29.39	18. 93	39. 41	24. 87	38, 02	1	•		- 1	
	PLACE4000670	7. 33	7. 03	6.77	5.23	9. 34	3.97	6, 41	3, 63	6.87				- 1	
	PLACE 4000685	27. 36	21. 25	28.71	7, 25	13.45	14.89	8, 25	13.79	8. 57		**	-	- 1	ı
	PLACE 4000687	5, 87	2, 72	5, 94	3.75	3, 52	3.04	2.8	3.66	2. 1	1			- 1	ı
	PLACE5000003		15.66	16.08	8.92	15.85	13.17	6, 36	9. 93	8.84		**		-	ı
	PLACE5000005	17.05	13.36	13.46	14.47	17. 35	13.76	5.61	8.29	7.64		**	ı	-	l
35	PLACE5000019			11.73	6.9	5.65	5. 55	3, 51	3.39	6. 14		**	-	-	ı
	PLACE5000021	28.4			18.99	19.21	16.52	23.2	21.13	21. 28	**	**	-	-	ı
	PLACE5000022		89.47	62.06	68.78	41.05	40, 85	49.9	41.64	44. 86 6. 39	1		ŀ		
	PLACE5800024			5.62	3.22	3. 37	3.62	2.52 80.48	4.61 55.18	59, 66	1		1		ı
	PLACE5000036			100.83	80. 5 5. 28	90. 12 9. 26	87.43 6.17	6, 41	11.67	8. 15	١.		۱	- 1	ı
40	PLACE5000059 PLACE5000076						121.21	73.46		129.94	1		1		ı
	PLACE5000117					20.06	15.63	5, 73	9, 53	5. 57			1	-	ı
	PLACE5000143					9.83	8, 06	5, 81	8. 66	8.8	1		1		ı
	PLACE5000152						116.95	74, 92	70.73	75. 92	1		1		ı
	PLACE5000154	90.81	108. 58			77.68	45. 59	40.84	23. 33	50.85			1	- 1	ı
	PLACE5000155	33. 2	24.58	26.07	16.94	31.52	23.72	16.67	17. 93	27. 02	l		1	1	l
45	PLACE5000165					103.03	88. 36	32.72			1	*		-	l
	SKNMC1000004			1083.3			€73.39				1		1		ı
	SKNMC1000011					12.47	5.45								ı
	SKNMC1000013					11.72	12.41	4.74				**	Į.	-	Ł
	SKNMC1000014					26. 13	18.1	14.76		11.78		**	1	-	ı
50	SKMMC1000018					9.73	4. 54	3.81 18.7	4. 41			**	1		ı
	SKNMC1000020 Sknmc1000046					25.33 22.27	12.68 16.07					••	1	-1	ı
	SKN#C1000046					4.73	4. 3						۱.		l
	SKMMC1000062					15.7							١-	1	١
	SKMIC1000075					10.78	7. 51						1		ı
	SKNMC1000073					13.04							1-		١
55	SKMMC1000091					23. 22						**	-	-	ı
	SKMMC1000099					7.5							1		١

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						aute 43	70							
	ISKNMC1000104	16.76	17.55	17.05	8. 52	14. 24	10. 52	9.72	10.59	13. 24	*	**	-	- 1
	SKNMC1000113	45. 53	34. 91	35, 51	20.7	25.14	23.85	18.86	24, 14	23.6	*	*	-	-
	SKNMC1000119	17, 58	14.67	21.54	10.33	12.31	8. 53	8.83	13, 02	8, 17	*	*	-	-
5	SKNMC1000142	16. 52	20.57	23, 49	7. 26	16.72	11.75	8.01	11.68	14.09		*		-1
	SKNWC1000170		183.41			226. 18		247.45	216.7	250.73		- 1		- 1
	SKNMC1000178	15.01	12.68	14, 32	6.9	9. 12	6, 49	4, 97	7.82	9. 01	**	**	-	- 1
	SKNMC1000194	7.33	8.61	11.61	2.23	3.73	3.19	2.64	4.74	3, 45		*	-	- [
	SKNMC1000198	18.76	29.87	24. 52	7.87	13.1	9.06	5.41	5.23	5. 02		**	_	-1
	SKMMC1000225		119.68	87. 32	52.94	84. 28	50.45	35, 91	33.78	44.06		**		-1
10		23. 35	22.68	30.9	13.07	19.76	17.19	12.83	14.11	14, 19				-1
	SKNMC1000249		10.7	16,74	6. 92	9, 27	7.23	6.86	7. 37	5. 89		: 1		- 1
	SPLEN1000007	10.7				13.65	10.56	13.22	11.71	11.02		٠,		- 1
	SPLEN1000012	15.31	13.75	18. 53	8. 26			29.87	29.67	37. 64		**	_	-1
	SPLEN1000014	74. 22	59, 99	63. 53	32.33	49.13	32. 58		3.76	2. 42	÷		-	-
	SPLEN1000036	14. 69	11.54	10.05	7.95	8. 19	6. 93	3.18	18, 35	17.18	÷	**	-	-1
15	SPLEN1000059	39,71	33, 5	32. 87	12.1	19.81	12.24	19.97			ï	**		-
	SPLEN1000068	20.63	22.57	21.07	11.6	16.29	10.46	9.19	11.89	8, 79			-	1
	SPLEN1000072	19.88	16.93	14.96	6.06	9.86	7.47	5.18	4, 49	6.78	**	**	_	- 1
	SPLEN1000101	71.97	43.16	57.46	18.69	27. 56	26.8	32.73	40.27	19. 97	*			- 1
	SPLEN 1000108	7.69	6. 28	7. 07	3.65	4. 35	2.67	3. 12	2.84	2.04	**	**	-	- [
	SPLEN1000113	11.47	6.64	12. 13	7, 54	8. 23	7.26	5. 5	6.27	7. 59				- 1
20	SPLEN1000114	16, 01	14.75	14.36	7.29	9.65	6. 3	8.14	8.22	8.73	**	**	-	- [
	SPLEN1000132	5. 96	4. 51	7.38	2.74	4.42	2.34	1.47	2. 25	2.21		*		-1
	SPLEN1000135	10.57	8.18	9. 39	3.86	4.71	4.73	4.9	2. 38	2. 97	**	**	-	- [
	SPLEN1000136	23.91	23. 93	20. 23	10.77	11.86	10.15	16.69	14.04	13.77		**	-	- [
	SPLEN1000141	42.88	39,73	42.83	27. 16	32. 91	23.4	23.31	18.41	20.57	**	**	-	- 1
	SPLEN1000164	15.72	13, 33	13.69	7.76	12.46	6.69	9.59	8, 11	9. 12		**		-
25	SPLEN 1000 166	12.98	9,08	11.45	4.56	6.01	5.04	6.49	5. 17	6.32	**	*	-	-1
	SPLEN1000175	19, 26	15.94	15.5	7.48	9, 23	6. 92	11.34	13, 49	13.41	**		~	-1
	SPLEN1000182	67.34	58, 35	68. 22	30,69	30.43	27.88	28. 16	29.1	23.7	**	**	-	-1
	SPLEN1000185	6.38	8, 35	5.56	4, 59	4, 56	3.06	2.86	2.31	1.51		**		-1
	THYMU1000004	48,79		41, 97	30.97	32.65	27.64	25.98	17.5	13. 12			i	- [
	THYMU1000009	14. 59		14.88	8,47	10.1	5	7.34	4.91	5.62		**	-	- 1
30	THYMU1000015	19.34		12.08	8.34	11.37	5. 93	10.39	10.96	13, 03			-	- 1
	THYMU1000016	6.78		5. 54	2.79	2. 51	1, 91	5.14	3.5	4.34	1		-	- 1
	THYMU1000023	6, 35		6.53	4, 39	3.65	4.2	3.1	5.69	3.98	**		-	- 1
	THYMU1000034			375.97		379.09	270,69	454, 49	361.08	359.56			i	- 1
	THYMU1000035	8, 93		9.14	5, 88	4. 95	4. 81	5.06	3, 93	4, 35	**	**	-	- 1
	THYMU1000037	9.69		7.36	4. 28	3. 96	3, 33	4, 24	3, 59	3.83	*	*	l -	-1
35	THYMU1000042	6.85		5. 82	2.75	3.24	2. 02	2.08	1, 63	1.09			-	- 1
	THYMU1000047	25.64		24.74	13, 43	16. 94	11.38	25.77	21.68	23.81	**		۱-	- 1
	THYMU1000080			27.22	17.42	19.98	26. 99	13.06	15.8	14.07			i	- 1
	THYMU1000094			9.51	6.45	6.49	4, 35	11.14	9, 41	9, 31	1		1	- 1
	THYMU1000109							235. 81	205.84				i	- 1
	THYMU1000127			37.5	12.05	18.11	17.75	24. 49	15.04	18.2	**		l -	-1
40	THYMU1000136			15. 84	7.65	12. 45	17.73	10.24	3.41	8. 22	1 *		-	-1
	THYMU1000137				3, 35	5. 2	3, 34	4.41	2.7				۱-	-1
	THYMU1000146				8. 11	7.24	6, 17	5. 13	3.03	6.19	1		i	- 1
	THYMU1000159			8. 17	6. 28	6.42	3, 77	4.54	3.62		۱.	**	l -	-1
	THYMU1000163			9.7	5. 33	5. 95	3, 66	7.57	4, 17			*	۱-	- 1
	THYMU1000167					31.06	17.27	75. 27	48, 55		1 **	•	۱.	- 1
45	THYMU1000186					4, 49	2, 65	3. 36	2. 91		1		1	- 1
	THYR01000017				8. 21	11.63	5, 25	6.04	5. 8		1		1	-1
						7.77	4. 55	4. 15	3. 45		1	i		-1
	THYR01000026					39. 02	28, 83	26.88	25. 91		i .	i	i	-1
	THYR01000034	38.0						22.09	19. 28		1	•		- 1
	THYR01000031				22.39	30.44	22. 35 4. 36	7.94			1		1	-1
50	THYR0100003					10.8			3.54			*	١.	-1
	THYR01000040						4 4 4	5.07				**	-	-1
	THYR0100006							6.42				**	1-	- 1
	THYR0100006							7.22					1~	
	THYR01000070											*	1	<u>†</u>
	THYR0100007					17.94		9.77				**	1	- 1
55	THYR0100008								8. 4				1	- 1
	THYR0100008											*	1-	- 1
	THYR0100008	13.5	6 10.42	8.08	7.5	9. 36	4. 87	6.64	6.5	4.47	1		1	- 1

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	THYR01000087	51.14	69.3	41. 47	29. 99	58.4	26.89	61.05	53, 08	61.77		- 1			
				30, 84						22.34		*	_	_ 1	
	THYR01000092	26. 07	28. 19		13.75	13.51	9. 94	17. 56	21.51		**	٠,		- 1	
	THYR01000093	4. 26	2.01	4. 16	1.77	3.46	3.25	2. 69	3. 3	2.41		- 1		- 1	
5	THYR01000099	9.62	5.87	9, 26	3.68	8.63	4. 69	5.9	7.16	4, 32		- 1			
														- 1	
	THYR01000107	70.48	59. 74	59.68	39.34	64.57	40.78	35. 76	61.82	50.34		- 1		- 1	
	THYR010001111	13.96	11.92	12.38	7.04	9.78	5. 89	9, 55	11.53	9, 66	*	*	-	- 1	
	THYR01000121	31, 45	32. 43	38.13	15,06	18, 17	17, 71	26.05	28. 32	34.79	**		_	- 1	
											**	- 1		- 1	
	THYR01000124	54.83	35. 35	48.22	37.6	35.37	31.52	34. 87	27.99	31.75		- 1		- 1	
	THYR01000129	8. 59	9.01	8, 49	5.14	9, 82	6.06	8. 93	8.73	10, 55		- 1		- 1	
10			8. 25		6. 62	6. 95	6. 32	6.99	9.76	9.68		- 1	_		
	THYR01000130	12. 23		10.08							•				
	THYR01000132	7, 93	8.79	121	5.35	7.15	7.04	4. 12	4, 66	3, 86		*		- 1	
	THYR01000134	17.2	6.06	9. 3	6.24	6, 27	12.5	5, 33	7.31	5.41		- 1		- 1	
								18.04	19.4	19.02		. 1		_ [	
	THYR01000144	23.66	30. 56	32. 27	27.55	32.97	28, 83					• 1		- 1	
	THYR01000155	35.26	37, 35	35. 3	18.99	24, 31	20, 42	31.31	40.6	29, 86	**	- 1	-	- 1	
	THYRO1000156	37.13	43.79	38.99	24.77	34.58	23.66	18.78	36.39	21, 81		- 1	-	- 1	
15										16.02	**	**	_	. 1	
	THYR01000163	56.18	53.4	58. 15	23.96	28.08	23.46	11.7	14.01		**		-	- 1	
	THYR01000173	8, 52	8, 35	6.58	5. 25	10, 18	4, 15	3, 25	4. 22	4,73		**		- 1	
	THYR01000186	35. 39	31.09	36.48	14.41	13.8	14.41	7.62	12.05	15.91	**	** ]	-	-1	
												**		- 1	
	THYR01000187	17.39	15.95	24.02	6.63	8. 73	8.87	5.49	7.02	6.9			-	- 1	
	THYR01000190	7.06	7.97	10.31	6.43	5. 1	6.9	4.78	5, 24	5.17		*		- 1	
20	THYR01000196	8.72	14, 15	13, 33	14.25	10.93	7,74	3, 75	6.65	9.84				- 1	
20														_	
	THYR01000197	13.75	17. 13	15.02	14.96	17. 34	16.5	8. 34	11.47	6.36		*		- 1	
	THYR01000199	6.8	9, 71	11.38	6.6	11, 32	10.32	5.5	6. 51	7. 17					
	THYR01000206	20.09	21.74	25.04	13, 14	17. 5	12.03	7.45	6.8	7.63		**	-	-	_
										3.66	٠.	*		_	
	THYR01000221	6.96	7.15	5.44	4.48	8. 31	3. 94	3.81	4. 45						
	THYR01000222	56, 33	56. 17	63.27	37.58	40.7	31.29	35, 13	33. 5	43.65	**	**	-	~	
25	THYR01000228		127.09	184. 25	73.36	64, 55	57. 59	55.06	55, 21	55.09	**	**	-	-	
20													_		
	THYR01000241	7.79	5.72	9. 15	4. 42	5	4. 49	2.68	4, 65	4.26	*		-	-1	ļ.
	THYR01000242	18, 16	23. 99	29. 76	25.09	51.85	27. 45	12.5	20.02	16.61					
	THYR01000246	21.57	18.12	24.56	19.46	20, 13	15. 33	9, 29	10.17	9.72		**		-	i .
	THYR01000253	10.77	13.3	16.4	9.6	16.01	8. 45	6.71	8.04	5.73		*		~	i i
	THYR01000270	12.58	12.99	17.24	8.11	13.02	8.87	9.06	8, 16	7.99		*		-	
30			13.53	16	8. 28	14.2	7.73	7.71	7.11	4, 92		**		- 1	i
50	THYR01000279											**			
	THYR01000285	1215.2	1221.3	1148.5	961.64	1121.8	816.57	1261.8		1518.7			1		
	THYR01000288	24, 11	22, 65	22, 01	13.89	25, 53	14.41	22.83	21, 93	20, 34					
								32.78	49. 02	40. 29				-	i
	THYR01000296		36.2	38.51	33.51	37.84	29.24				١.		1		i
	THYR01000320	12.29	9.4	14.15	6.59	7.17	6.08	5.28	11.64	9. 29	*		-		
	THYR01000322		34.76	35.08	17.69	26.05	19.02	19.43	23.34	20.74	**	**	-	-	į.
35						27. 29		10.07	12.24	13.52			ł	-	
	THYR01000327		26.35	19.48	16.99		15.75				١.		1		
	THYR01000343	16.65	12.58	17.82	8, 18	11, 42	7.21	10.1	9. 05	8.44	*	*	١-	-	1
	THYR01000345	6.25	7.77	6.06	4.74	8.59	3.02	4, 68	4,72	4, 12			1	-	
	TITTRO 1000040		10.33	11.35	6. 24	8.76	7.32	8, 15	6.51	9.21	۱.	*		-	1
	THYR01000358											•	١.	- 1	
	THYR01000368	5.38	5. 26	6. 26	3.15	3. 94	2.68	6.19	3.06	2.43	**		-	- 1	
	THYR01000375	14.1	11.6	18.84	8, 56	5. 99	5.1	3.89	3.2	3, 55	i *	**	l -	-	ł
40	THYR01000381	10.31	8. 3	9.78	4.37	5.3	5.96	4, 17	5. 35	3.59	**	**	-	-	
													l		
	THYR01000387	11.39	10.61	13.54	4.54	5. 97	3. 46	10.59	5.84	3. 07	**		1 -		ı
	THYR01000394	61.78	50. 97	22.02	14.67	26. 91	8, 08	6.98	9.73	11.33			1	- 1	ł.
	THYR01000395		37.87	27.54	18.13	24. 45	14, 15	25. 79	23.19	25.7	۱.		۱.		
										3.08	1	**	1	-	
	THYR01000400	10.51	10. 51	9. 69	3.23	12.01	5.83	3,72	3.73		1		1		
	THYR01000401	10.42	9. 53	10.77	6.74	5, 06	4.73	3.24	5, 18	5. 6	**	**	<b>!</b> -	-	
45	THYR01000407		92.99	98.17	63.29	75.05	58, 17	51.33	55, 33	55.02	1 **	**	۱-		
												• • •	١		
	THYR01000420		13.25	17. 92	11.1	11.62	8. 56	12.79	12. 37	14.41	١.		1 ~		
	THYR01000438	8.38	5.22	8.85	5.12	4.11	4, 31	4.22	4.7	3.13	1		1	-	
	THYR01000452		12.59	12.97	5.92	10.6	6. 13	2, 16	3.93	4.57	1 *	**	I	-	ı
											1 **		-	-	
	THYR01000455	7.17	8, 57	9.52	2.62	3 31	3.44		2. 18	2. 11	1 **	**	١-	-	
	THYR01000471	78.16	81.99	62.03	65.6	82. 48	70.86	81.77	51.28	74.06			1		
50				8. 46	4.91	7.54				5.02	1	**	1	-	
	THYR01000481										١.				
	THYR01000484		12.76		10.37	11. 59			9.7	11.91	۱ *		١~		
	THYR01000488	28.64	20.75	23. 35	20.89	21.13	19.59	15.6	20.62	19.42	ı		1		
	THYRO1000501				7.74	9, 03	6.79			10.26	1 ##	*	I -	-	
											1 *		-	_	
	THYR01000502				44.07	46.78		44.58		38. 11			1 -	-	
55	THYR0100050	9,64	7.65	7.27	5.56	5. 82	3.06	3.98	5. 47	2. 02			1 -	-	
w					38.72	38. 24				18.32		**	1	-	
	THYR0100053	7 . 31 . 32						1	EV. 31			**	1	_	
	THYRO1000558	il 102. 92	92, 57	85.3	76.31	104. 96	62.73	51.56	50.77	49.93	1	**		-	

					7	able 5	00							
	THYR01000558	8, 54	7.17	6.38	4, 15	8.61	3.01	4. 25	2.04	3.45		*		-1
	THYR01000569	6. 63	4. 92	4.79	4.77	4	3. 41	3. 56	3.02	3.94		*		-1
	THYR01000570	9.09	6.95	8.33	5. 27	6.38	8.54	5.25	4.89	7.09		- 1		- 1
5	THYR01000572	52.16	31.52	39.78	12.92	33.15	30.73	30.68	23.84	11.23		- 1		- 1
	THYR01000573	8.96	8. 49	6.82	4. 33	8.49	6.35	13.45	11.85	8.38				- 1
	THYR01000577	9.73	7.83	7.69	3.93	6.26	2.6	2. 58	3.82	2.74	*	**	-	~
	THYR01000580	40.56	32.54	29.95	22.22	19.86	19.81	24. 19	17.34	25. 52	*	*	_	- [
	THYR01000584	39.96	36.8	31.93	19.25	25. 5	16.24	26. 15	21.02	19. 13	*	:	-	-1
10	THYR01000585	8.99	7.37	6.65	4. 19	6.67	4.86	5. 4	3.58	2.99		*		-1
	THYR01000596	43.85	33.14	32.61	22. 94	31.65	11.59	15.52 8.87	10.88	14. 47		1		- 1
	THYR01000602	19.04	15. 25 9. 44	10.78	7. 23 5. 29	11.51	11.78	4.9	5. 21	5, 08		i		- 1
	THYR01000605	10.06 134.15	117.37	5. 91 85. 97	86, 81	98. 31	67. 18	77,31	67	84. 37		- 1		- 1
	THYR01000615	33, 27	24. 69	16.68	12.6	10.75	20. 13	8.38	6, 85	11.76		.		-1
	THYR01000636	17.91	16. 49	14. 48	12.48	11.49	6.7	8.72	6. 17	8, 59	*	**	-	- 1
15	THYRO1000637	18. 22	22.2	13.43	10.39	14.05	9.78	9, 07	5. 2	7.26		*		- 1
	THYR01000641	7.23	5.87	6. 32	4. 82	10.06	3.87	8, 42	4.55	2. 07		-		١
	THYR01000657	10.91	9.86	8.65	6. 93	7.74	7,68	9. 53	8.35	10.94	*	- 1	-	- 1
	THYR01000658	14.37	9, 48	5.1	6. 28	6. 83	5.43	4, 59	4.62	4.77		- 1		- 1
	THYR01000662	10.63	9.43	8.46	6.7	6.85	6.96	7. 92	7.09	10.89	*	- 1	-	- 1
20	THYR01000566	30.51	33.5	45. 43	12.9	20. 33	13.81	14. 6	13.53	16.48	*	**	-	-1
	THYR01000676	11.56	11.35	8. 48	6.32	9.3	8. 37	4. 16	5.02	6,44		*		-
	THYR01000678	23.44	18.73	20.73	10.13	15. 99	8. 49	. 9	7.38	9. 87	*	**	-	-
	THYR01000684	27.78	27.85	20. 27	12. 26	16. 45	11.06	26.97	17. 92 9. 36	24.98	*	- 1	-	
	THYR01000694	16.87	11.78	10. 72	5. 47	10. 25	5. 99 3. 48	12.29 7.37	3.08	8. 6 5. 06				1
	THYR01000699	9. 11 17. 55	6. 35 14. 87	4. 55 12. 31	5, 62 9, 08	6.31 11.99	7.57	12.79	9. 26	13		- 1		1
25	THYR01000712 THYR01000715	29.82	23. 25	21.02	20. 98	27. 37	20.44	15.76	18. 98	18. 68		1		1
	THYRO1000716	13. 12	15, 14	11.94	8.87	12	7.65	7.48	6.69	5. 98		**		
	THYR01000717	11.89	15. 21	12, 45	6.9	9. 22	5.08	30: 93	6.96	7, 18	*		-	- 1
	THYR01000723	30, 36	26.4	21.78	10.82	20, 63	14, 48	30, 94	22. 3	30. 26			-	
	THYR01000734	16.61	19.91	16. 12	11. 94	17.61	9, 39	9, 31	6.61	13.3		*		-
30	THYR01000748	17.46	14.06	15.87	5	7.52	5.42	7.82	5.84	7.09	**	**	-	-
	THYR01000755	20, 17	21, 24	19.88	13.28	17.61	14.04	14.7	23.04	21.01			-	
	THYR01000756	369.81		369.63	298. 51	345.41		259.37	366.07	273.4				1
	THYR01000776	4.97	4. 99	7, 55	4.25	6. 18	4. 82	2. 58	5.02	3. 55				
	THYR01000777	5. 42		6.34	5.34	6.01	4.03	3.82	3.75	3. 21		**		-
	THYR01000779		458.9	483.51		549. 42	449. 1	368.6	510. 93 15. 4	458. 47 16. 35				
35	THYR01000782	17.27 5.74		23.35 6.68	14.58 5.43	17. 12 9. 62	13.37	15.61	5. 19	3.8				-
	THYR01000783 THYR01000786	25.62		26.26	16.78	13, 18	14.48	32.89	30.78	25.94	**	•	-	- 1
	THYR01000787	50.52		54.05	28.71	18.06	26. 25	20, 11	24.5	22.55		**	۱-	-
	THYR01000792	14.11		15.65	4. 55	6.72	7.1	8, 13	7, 14	8. 33	**	**	-	-
	THYR01000793	36, 92		42. 61	19.08	27. 16	19.55	34, 91	38.74	32.64	**		~	
40	THYR01000795	37. 33	45, 29	88.32	42.19	57.36	40.15	12.57	21.17	13.38			l	
	THYR01000796	10.25		14.01	6.19	10. 21	5. 98	7	11.89	11.1				
	THYR01000798	9. 87		12. 83	9.06	8. 1	7.49	5.04	7.33	5, 2	*	*	-	-
	THYR01000800	37.69		19.73	33.2	22. 36	23. 21	13.79	16.89	13.75				
	THYR01000805	8. 27		6.02	5.33	8. 39	4.44	3.81	4. 14 21, 16	3.82 31.98	**	*		_
45	THYR01000815	109.41		90.62	43.51	38.14	25.61 18.45	31.76 15.87	14.34	12.65	*	::	-	-
40	THYR01000829 THYR01000835	33.86 9.51		34.71 10.3	19.68 6.71	21.68 6.28	4, 65	5, 41	5. 24	6. 07	**		1 -	_
	THYR01000835	27.06		24, 96	16.28	19.34		6.22	7.81	8. 06	**	**	1	_
	THYR01000846	7.86		13.48	4.53	6. 28	7.74		7.02	5. 26			ı	
	THYRO1000852	24.65		23.66	12.87	20. 2	12. 5	9.33	10, 39	11.62		**	-	-
	THYR01000855			10. 43	2.52	5, 79				5. 2			-	
50	THYR01000865			8.64	3.4	6, 56	3, 18	5, 99	8.76	7.67			1	
	THYR01000866	57.27		51, 69	32.85	43, 74	27.97	46.24	39.05	41.55	*		-	
	THYR01000881	13.67		13.18	9.6	10, 62	12. 24	10.21	16.82	10.91			1	
	THYR01000894			11.4	5.66	6.12	4. 31	4.14		4.75		*	-	-
	THYRO1000895			10.11	3.35	4.07	4.72			3. 45		*	١-	-
55	THYRD1000916	9.11		16.87	5.02	6.67	5. 27	4.15	6. 12	3. 42	1			
55	THYR01000917			12.32	4.97	9, 51			7.94	6.2			1	
	THYR01000926	14.99	10.81	12.37	6.79	9, 95	5.06	7.42	7.03	11.27			1 -	

THYROLOOPSI   1.19   1.38   1.72   1.75						Т	able 50	01								
THYROLOUGUSS   St. 82		THYR01000934	9.21	6.94	9. 67	4. 17	7.52	4. 261	6,41	5.73			*		-1	
THYRD		THYR01000951	21.19	13.82		10.15	10.71	6.54	7.11	11.34						
		THYR01000952	38. 82	45.17	51.97	24. 89	24.37	18.77	17.55					-	-	
THYRRO1000981   31,04   51,28   36,49   15,01   15,48   15,21   21,6   24,41   21,14   32,46   17,17   31,92   67,7   51,62   31,7   41,7   42,7   41,7   41,7   42,7   41,7   41,7   42,7   41,7   41,7   42,7   41,7   41,7   42,7   41,7   41,7   42,7   41,7   41,7   42,7   41,7   41,7   42,7   41,7	5	THYRO 1000956	11.42	8.41	13.85	5. 27						*	*	-	- [	
THYRDIOLOGY   14.43   12.46   17.93   1.92   16.7   5.25   5.24   17.53   4.58   22   23   24   25   25   25   25   25   25   25		THYR0 1000960													-	
THYRRIODOST														-		
THYROLOGOST   7.51   5.29   9.43   2.24   6.94   1.8   3.55   5.95   3.45   1.14701000951   1.14701000952   3.5   4.14   7.44   3.85   5.01   3.16   3.1   3.1   4.71   4.75   4.75   4.75   3.15   4.75														-		
THYROLOGOSIS   7.71												*	**	-	-	
THYROLOOSS   S. 4   A. 14   7.44   S. 85   S. 03   3.16   3.11   3.12   4.03	10							1.8					- 1		-	
THYROLOUSSAI   S. 35													- 1		- 1	
THYROIODSSS   10.2   6.57   7.52   4.27   4.23   7.46   2.54   2.55   2.56   2.52   3   48   -														_	_	
THYROLOUGUS   12.33   10.95   12.47   5.27   7.49   7.34   5.57   6.15   6.10   12.35   1.75   7.7														_		
THYROIODISSS   9.77   12.44   10.44   5.68   8.33   5.29   5.68   3.11   2.9																
THYROLOUDIC   1.65   1.37   1.16   7.36   1.38   1.48   1.59   7.17   2   509,02   394.65   380,17     THYROLOUDIC   1.65   1.37   1.16   7.36   1.38   1.48   1.77   1.05   27.79   31.88   30.58   27     THYROLOUDIC   1.575   5.97   11.29   5.12   7.05   5.98   6.11   3.9   3.44   2   2     THYROLOUDIC   1.575   5.97   11.29   5.12   7.05   5.98   6.11   3.8   3.05   2   2     THYROLOUDIC   1.34   7.37   5.88   7.55   5.52   6.01   4.3   4.39   3.44   2   2     THYROLOUDIC   1.34   7.37   5.88   7.55   5.52   6.01   4.3   4.39   3.44   2   2     THYROLOUDIC   1.38   3.8   6.8   5.14   7.7   1.58   5.55   4.32   2.78   4.8   2     THYROLOUDIC   1.38   3.8   6.8   5.14   7.7   4.58   3.55   4.32   2.78   4.8   2     THYROLOUDIC   1.58   5.7   5.75   5.88   7.55   5.8   5.55   4.32   2.78   4.8   2     THYROLOUDIC   1.18   4.7   5.7   5.98   4.77   3.1   5.00   4.4   4   4   4   4   4   4   4     THYROLOUDIC   1.18   4.7   5.7   5.98   4.77   3.1   5.00   4.4   4   4   4   4   4   4   4   4														_	- 1	
THYROLOUIDIS   6.58   1.37   1.65   7.36   13.81   8.4   12.04   9.04   10.47	15														- 1	
THYROLOUID   34.7   29.76   12.31   14.48   17.7   10.51   27.97   31.88   30.88   27   11.77   10.78   11.77   11.7													-		- 1	
THYROLOGICAL   1.57   5.59   5.71   7.72   20.81   6.88   8.19   10.85   5.15   6.58   8.8   8.8												**	- 1	-	- 1	
THYROID0131   15.75   5.77   1.28   5.12   7.05   5.99   8.11   6.39   3.44   2   -										6.58	6.8	*	**	-	-	
THYRDIODICAL   1.3											9.34		- 1	-	- 1	
THYRRIDOIDED   12.86   12.45   12.49   4.88   5.59   4.88   1.55   4.32   2.35   * * * *	20									4.01	4.9				- 1	
THYROLOGICAL   1.0, 18, 18, 18, 10, 18, 18, 18, 18, 18, 18, 18, 18, 18, 18							5, 59	4.88	3, 54	3.68		**				
THYRRID   100   100   1.77   1.2 0.6   1.47   3.65   1.2.27   3.17   3.06   4.59   4.41   \$\$ = \$\$ - \$\$		THYR01001063														
THYROLOGIUS   1.2, 17   5.3   6.75   5.82   4.91   5.8   5.85   8.31   7.24   12   14   14   14   15   15   15   15   15												**		-		
**   **   **   **   **   **   **   *													**		-1	
THYROLOGIUS   0.18   5.7   5.85   5.04   6.9   4.52   4.18   4.71   4.45   1.71   4.75   1.												**	. 1	-	. 1	
THYROLOGIUS   S.1   Z.75   Z.78   0.89   0.92   1.62   1.22   0.94   0.7   * * *   *   -	25												*		1	
THYROLOUILOR   S. S. L. L. S. O.   4., 26   10., 38													. 1		- 1	
THYROLOGIUS   0.89   5.97   7.89   2.84   3.69   2.77   2.56   1.4   2.3   4   22   4   4   5   5   5   5   5   5   5   5														-		
THYROLOGIZE  20.44   16.36   16.59   19.11   13.05   6.39   13.27   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   2   15.75   13.77   15.75																
THYROLOGIS   20.41   6.36   6.91   9.1   12.05   6.92   13.22   15.3   13.97   2   -																
THYROLOGITA   25.00   18.24   17.74   2.83   2.81   2.74   2.84   2.77   1.82   2.85   2.81   1.85   2.85	20												"	_	- 1	
THYROLOUIS 7, 42 5.4 5.48 4 5.68 4 5.50 4.22 3.88 4.14 3.81 * * * HYROLOUIS 7, 42 5.22 3.68 5.22 8 1.65 12 0.55 18.49 2.21 6 16.92 2.07 * * * HYROLOUIS 7, 42 5.50 4.99 5.1 2.78 3.05 3.47 2.44 2.26 1.6 * * * HYROLOUIS 7, 18.8 5.64 2.94 2.03 6 1.47 2.44 2.26 1.6 * * * HYROLOUIS 7, 5.55 8.04 8.05 1.17 1.18 5.10 4.7 1.18 5.15 3.3 1.18 1.7 * * * * HYROLOUIS 7, 5.55 8.04 8.05 1.17 1.18 5.10 4.7 1.18 5.15 3.3 1.18 1.18 * * HYROLOUIS 7, 5.55 8.04 8.05 1.17 1.18 5.10 4.7 1.18 5.15 3.3 1.18 1.18 1.18 1.18 1.18 1.18 1.18	30												**	_	-	
THYROLOGITISA   28,23   21,26   23,28   6,51   20,53   18,49   22,14   16,52   20,47   8   -													**	-	- 1	
THYROLOUIT   1.8   5.04   4.99   5.1   2.78   2.05   3.47   2.44   2.26   1.6   32   43   3.57   3.79   3											20. 47	*		-	- 1	
THYROLOUITS   31.85   26.04   28.94   20.36   18.72   13.59   18.5   16.5   15.33   18.17   2 **								3.47			1.6			-		
THYROLOUITS   S.5.55   S8.04   80.56   10.29   11.41   8   77.1   51.33   46.56   49.17   84.5   - 1				26.04	28.94	20.36	18, 72	13.59	18.5			*		-		
Intrincipolity   10.9   11.18   11.71   5.13   7.17   5.9   4.63   5.82   5.02   ** **   -     Intrincipolity   11.83   6.2   11.19   5.29   7.22   4.28   5.84   7.01   6.85   6.85     Intrincipolity   11.83   6.2   11.19   5.29   7.22   4.28   5.84   7.01   6.85   6.85     Intrincipolity   11.83   6.2   11.19   5.29   7.22   4.28   5.84   7.01   6.85   6.85     Intrincipolity   1.64   3.14   3.65   5.29   7.22   4.28   5.84   7.00   6.85   6.85   6.85     Intrincipolity   1.64   3.14   3.65   7.28   7.85   7.85   7.84   4.82   **   -     Intrincipolity   1.64   3.14   3.65   7.28   7.85   7.85   7.85   7.85   7.85     Intrincipolity   1.67   7.02   5.87   1.68   7.85   7.85   7.85   7.85     Intrincipolity   2.7   2.7   2.85   2.85   2.85   7.85   7.85   7.85     Intrincipolity   2.7   2.7   2.85   2.85   2.85   7.85   7.85   7.85     Intrincipolity   2.7   2.7   2.7   2.85   2.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85   7.85     Intrincipolity   3.85   3.85   3.85   3.85   3.85   3.85   3.85   3.85   3.85   3.85     Intrincipolity   3.85   3.85   3.85   3.85   3.85   3.85   3.85   3.85   3.85     Intrincipolity   3.85	35	THYR01001175		98.04	80, 56	104. 29	114.18									
THYRRIDOIDIS4   11.83   6.2   11.19   5.29   7.22   4.28   5.84   7.01   6.86   1.76			10.9	11.18								**	**	-	- 1	
### THYROUGIZEM \$2, 04 31, 32 2, 32, 81 8, 22 19 8, 22 7 19, 75 37, 94 46, 59 \$ - 1		THYR01001189													- 1	
THYRRODICIZES   8.98   12.22   9.39   5.39   8.08   5.04   6.55   6.04   4.85   8   0   - 1															- 1	
### HTMF01001213 41.54 37.72 35.56 23.56 25.7 20.79 37.67 32.41 38.62 4** - HTMF01001224 10.56 11.43 10.75 5.38 6.96 5.7 7.21 4.53 6.56 ** * - HTMF01001224 11.57 62.25 15.97 16.59 12.69 5.38 9.76 5.69 5.27 4** - HTMF01001225 12.1 19.03 16.06 12.27 6.61 5.71 2.27 10.55 68 5.27 4** - HTMF01001253 27.32 21.68 20.65 87.4 18.6 16.73 19.21 5.95 9.02 8 8 HTMF01001253 27.32 21.68 20.65 87.4 18.7 18.7 17.7 10.5 18.6 13.04 8 8 8 1.04 8 8 8 1.04 18.7 18.7 18.7 18.7 18.7 18.7 18.7 18.7		THYR01001204								37.94	40. 39			-	- 1	
	40												•		~ [	
THYROLOGIZAT   16,776 20,25   15,72   16,59   12,69   5,38   9,78   5,59   6,27   22   -     THYROLOGIZAS   12,1   19,01   16,06   12,27   6,61   6,77   6,61   6,77   6,27   1,59   6,00   4   2   -     THYROLOGIZAS   27,12   21,58   20,56   8,74   18,76   12,77   10,51   6,56   13,04   4   2   2     THYROLOGIZAS   25,46   17,38   17,48   9,56   11,28   8,76   13,97   15,24   19,55   4     THYROLOGIZAS   25,46   17,38   17,48   9,56   11,28   8,76   13,97   15,24   19,55   4     THYROLOGIZAS   126,74   12,77   14, 12,77   14, 13,77   12,52   18,92   22,105     THYROLOGIZAS   126,74   126,77   13,88   4,97   7,48   6,55   4,02   44,54   8,97   22,105     THYROLOGIZAS   13,9   15,44   13,22   8,55   8,53   8,57   8,50   8,54   4,27   7,5   8,50     THYROLOGIZAS   13,77   12,77   11,0   6,16   6,14   4,27   4,57   5,3   8,04   4,27   4,57   5,3   8,04     THYROLOGIZAS   13,77   12,77   11,0   10,77   13,77	40															
THYROLOUIZE   13.1   18.0   16.05   12.27   8.51   6.73   9.21   5.95   9.02   \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$												٠.				
THYROLOUISS   27, 32   21, 58   20, 55   57, 41   8.78   12, 77   10, 51   6.86   13, 04   \$ **   -														-		
THYROUOIZEZ 8.57 4.74 4.05 1.76 5.55 4.02 4.79 1.39 5.05   THYROUOIZEZ 5.45 17.38 17.48 1.95 17.28 8.76 11.37 15.24 1.95 5.8   THYROUOIZEZ 5.45 17.38 17.48 1.95 17.28 8.76 11.37 15.24 1.95 5.8   THYROUOIZEZ 17.85 25.6 17.14 13.77 18.43 9.74 12.54 18.8 22.30 5   THYROUOIZEZ 18.67 17.55 18.27 15.76 18.60 71.99 10.18 93.81 120.13   THYROUOIZEZ 11.9 16.44 13.22 8.55 18.65 18.65 18.5 2.20 5 6 8 9 -   THYROUOIZEZ 11.9 16.44 13.22 8.55 18.3 18.9 5.3 1.64 6.6 9 9 -   THYROUOIZEZ 18.17 17.01 9.99 19.11 8.25 41.39 28.42 18.66 9 9 9 -   THYROUOIZEZ 28.89 24.47 17.01 9.99 19.11 8.25 41.39 28.42 18.6 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9												1	**	-	- 1	
45   THYROLOUISE   25.45   17.30   17.45   8.95   11.25   8.76   11.97   15.24   19.55   8   -   THYROLOUISE   72.75   25.6   17.45   13.77   14.1   13.77   13.41   3.74   12.55   18.92   23.05   THYROLOUISE   11.2   8.72   8.52   8.57   16.96   71.99   100.16   93.81   120.13   THYROLOUISE   11.3   8.12   8.56   4.57   7.48   6.55   1.36   5.44   8.4   8   -   THYROLOUISE   13.9   15.44   13.22   8.55   8.33   1.9   8.33   1.36   5.44   8.4   8   -   THYROLOUISE   13.77   12.27   11.41   6.73   8.58   4.22   7.87   7.87   5.3   8.64   8.9   8.9   THYROLOUISE   13.77   12.27   11.41   6.73   8.58   4.22   7.87   5.3   8.64   8.9   8.9   THYROLOUISE   13.77   12.27   11.41   6.73   8.95   1.27   12.57   12.97   2.22   1.98   THYROLOUISE   4.58   10.44   8.95   8.95   8.95   8.57   8.67   8.67   8.67   8.58   8.23   7.97   1.25   THYROLOUISE   15.07   13.59   11.8   5.59   4.89   8.67   80.61   56.13   42.33   8   9   9   THYROLOUISE   20.70   13.95   13.64   13.81   13												1			- 1	
THYROLOGIZII 27, 85 25, 6 17, 14   13,77 18,43 9,74   22,54 18,52 23,05   THYROLOGIZII 12,76 17,76 18,77 18,43 9,74   22,54 18,52 23,05   THYROLOGIZII 13,9 18,44 13,22 8,55 8,49 7,7,48 6,55 1,56 5,54 8,41 9 7 14780101297 13,17 12,27 11,41 8,22 8,55 13,23 3,9 8,23 3,56 8,23 3,56 6,5 8 9 1,77 14780101297 13,17 12,27 11,41 8,73 6,89 4,21 7,87 5,3 8,04 8 5 7 147801001313 3,4 2,28 2,88 2,47 17,01 9,39 19,11 8,25 4,139 28,42 3,56 8 9 1,47 17,10 19,10 1,47 1,47 1,47 1,47 1,47 1,47 1,47 1,47	45								13, 97	15.24	19.55			-	- 1	
THYROIODIZET  128,74   172,53   28,71   56,76   169,06   71,99   102,16   93,81   120,13   177,17										18.92	23.05			1	- 1	
THYROUOIZET   13,9   15,44   13,22   8,85   8,33   3,9   8,83   3,84   6,66   8 ***   -				172.63			169,06	71.99	103.16			1			- 1	
THYROUOLIZIN 13. 17 12. 27 11.41 6.73 6.88 4.23 7.87 6.3 8.04 82 22   THYROUOLIZIN 28.98 24.47 17.01 9.99 19.11 8.25 41.39 28.42 13.69 28.47 13.69 17.47 17.01 9.99 19.11 8.25 41.39 28.42 13.69 28.47 13.69 17.47 1		THYR01001290	11.3	8. 82	9,68	4.97	7, 48							l -	- 1	
THYRO1001302   22.98   24.47   17.01   9.99   18.11   8.25   41.39   28.42   18.98		THYR01001291	13.9	16.44	13.22	8.85								- 1		
INTRODUCTION   26.386 2.44   17.54   3.58   3.58   3.57   3.57   3.58				12. 27								**	**	-	- 1	
THYRO1001320 4,85 2.3 4.45 18.295 2.21 2.57 2.22 1.94 THYRO1001321 84,54 108.43 88.51 45.59 48.9 36.67 60.61 55.13 42.33 45 5 - THYRO1001327 15.07 13.59 11.8 5.6 6.9 4.45 5.91 6.73 6.71 45 5 5 - THYRO1001327 61.14 62.79 47.55 27.21 38.08 29.25 40.32 44.88 30.36 45 5 - THYRO1001327 21.04 18.95 13.54 17.83 15.44 11.88 17.21 14.24 12.55	50	THYR01001302	28.98							28. 42		١.		l	- 1	
THYROUOUSZI 44.54 108.42 89.51 45.59 48.9 31.6,77 60.61 56.13 42.33 45 9 - THYROUOUSZI 51.07 13.59 11.8 5.69 4.45 5.9 4.45 5.9 4.75 12.71 45.71		THYR01001313	3.4									۱*		-	- 1	
THYRO1001322 15.07 13.59 11.8 5.6 5.9 4.45 5.91 6.73 6.71 ** ** THYRO1001327 61.46 62.79 47.56 27.21 36.08 29.25 40.32 44.88 30.36 ** * THYRO1001318 21.04 18.95 13.64 17.33 15.44 11.88 17.21 14.24 12.15												١.,	_	١	- 1	
THYRO1001327 61.14 62.79 47.56 27.21 36.08 29.25 40.32 44.88 30.36 ** * THYRO1001336 21.04 18.95 13.64 17.33 15.44 11.88 17.21 14.24 12.35														1 -		
55 THYR01001336 21.04 18.95 13.64 17.33 15.44 11.88 17.21 14.24 12.35														I.		
	55												7	1	1	
													**	1	-	
		THYR01001347	42.47	35. 41	, 29.50	20.44	31.43	14.1	1 20.21					•		

					-	Table 5	02							
	THYR01001358	6, 96	7.77	6, 52	4. 41	8.39	3.78	5. 37	5. 5	4. 22		* 1		- 1
	THYR01001363	36.77	35.26	36. 97	12.86	20, 63	14, 65	24, 19	31.52	25. 12	**	*	-	- 1
	THYR01001365	2.39	1.31	4. 42	2.3	2.69	1.43	1. 82	2.18	1.5				ı
5	THYR01001374	10, 11	11.14	11.66	5. 19	7.75	3.64	2.84	5. 3	4. 44	*	**	-	-
	THYR01001401	11.45	8.59	8.77	5.02	5.3	5. 96	5. 5	8.56	6. 31	*	- 1	-	- 1
	THYR01001403	3.12	3.62	7.98	1.61	2.43	2.61	3.73	1.98	3.91		- 1		- 1
	THYR01001405	84.82	60.76	92.35	68. 75	78.6	50.19	53.53	29.6	47.94		*		- 1
	THYR01001406	13.59	7. 57	12.02	4. 76	6.87	3.76	11.59	13.59	9. 54	*	- 1	-	- 1
10	THYR01001411	4.72	4. 15	5. 65	3.88	2.4	3. 38	2.52	2. 5	3.6		*		- 1
10	THYR01001420	9.34	4. 82	8.71	4.7	4.71	5. 55	3.24	4. 39	8.81		- 1		- 1
	THYR01001426	9, 26	4. 52	12. 27	5. 65	6.22	4.37	4. 27	8. 59	6. 13		- 1		ı
	THYR01001430	10.74	13. 91	18. 18	13. 99	18. 28	13.49	6.82	9. 48	10.77		- 1		- 1
	THYR01001434	7.66	6.66	12.91	8. 64	10.68	8. 13	5. 38	8. 91	2. 99		- 1		- 1
	THYR01001456	26.49	36.99	36. 96	17.02	23. 99	16. 92	30.71	34.81	38. 85	*		-	- 1
15	THYR01001457	14.33	10.91	13. 26	5. 72	4. 54	3. 33	4, 05	3. 22	4.83	**	**	_	-
	THYR01001458	15.32	13.83	12.78	10. 99 2. 81	10. 13 3. 46	5.72 4.24	4.4	7. 29 4. 43	5. 51	÷	**	-	~ [
	THYR01001459	5.78 13.48	10.81	7. 36	5. 13	9. 62	6. 19	3.11 6.36	8. 85	9. 45	÷		_	-1
	THYR01001471 THYR01001478	8.04	3.79	8.01	2, 94	3. 62	2. 21	4.12	4. 22	3. 59	٠	•	-	- 1
	THYR01001480	16.07	16.86	14, 48	6.78	13.68	13.34	3.54	6.61	19. 35				- 1
20	THYR01001481	16.23	13.97	17. 19	13.3	15. 2	13.73	4. 94	6.01	3.76		**		-1
20	THYR01001487	39. 34	43.94	45.87	41.49	42.62	34.09	36.78	39.23	41.44				- 1
	THYR01001495	6, 21	3.33	12.71	2, 18	5. 51	4. 67	3.43	3.46	4.85				- 1
	THYR01001498	13,82	15, 45	16.61	11.32	20.15	8. 37	4.81	6, 08	4. 31	-	**		- 1
	THYRO1001510	9, 97	6.87	19. 31	8.41	6, 21	3.55	4.99	6.45	19. 57				1
	THYR01001512	11.51	8.21	10.34	5. 87	6. 36	6.84	4.43	19.67	5. 22	*		-	- 1
25	THYR01001519	12.05	16.47	10.9	4.73	8.69	4.14	4.42	4.28	5.04	*	**	-	- 1
	THYR01001522	10.39	6.84	12. 83	6.15	7.71	8	5.13	5. 33	7.56				- 1
	THYR01001523	15.14	10,51	16.01	7.01	9. 2	6. 38	3.77	3.19	4.5	*	**	-	- 1
	THYR01001526	5.67	5.48	12.84	2.85	5.19	3.09	4:51	3.63	4.01				- 1
	THYR01001529	70, 52	99. 68	61.07	73.62	81.82	35. 88	29.07	23. 9	34.69		*		-1
	THYR01001534	10.48	7. 92	11.72	6.09	7.42	11.24	6.53	6.35	3.17		*		-
30	THYR01001537	5. 9	3.05	5. 93	3.4	3. 75	2.75	3.25	2.87	4. 91				- 1
	THYR01001541	27. 84 12. 87	38.72 9.76	38. 42 11. 61	23. 51 3. 56	30. 22 6. 9	18. 99 5. 41	22.01 4.15	26.98 22.14	14. 85	**		_	- 1
	THYR01001545	143.67		127.43	126. 92	211, 99	217.97	165.83	143. 91	101.7	**		_	- 1
	THYR01001563	29.66	25. 88	28. 83	14.46	17.77	19, 17	10.82	18.32	17.35	**	**	_	- 1
	THYR01001570	5.85	5. 24	10.72	3. 21	3.79	2. 31	2.47	4.03	3.05	**			
35	THYR01001573	10. 58	14.52	15. 67	7.66	10.98	10.26	4, 93	9.17	5.76				-
	THYR01001584	11.71	9.79	13.05	7.75	12.43	7.3	8, 19	4. 92	4.88				- 1
	THYR01001593		87.93	122.57	37.27	54. 3	33, 56	100, 89	84.41	119.61	**		-	- 1
	THYR01001595	9, 11	9. 13	10.8	7,84	8, 88	6,77	4.62	12.97	7.82				- 1
	THYR01001596	6.25	3.54	8. 97	3.03	4. 5	2.76	3.54	3,67	4.02				- 1
	THYR01001602	7.1	6.21	7.8	6, 63	4. 96	5. 02	4.21	4.12	2.97		**		-
40	THYR01001605	28. 22	39.54	25. 87	13.48	23.89	22.57	11.83	10.02	13.09				-
	THYR01001608	19. 26	17.94	17.77	8.3	22	8. 49	8.07	8.36	11.18	١.	**		-
	THYR01001617	7, 36	9.07	12. 26	4.4	5, 61	3, 53	3, 83	3.37	4. 51	*	*	-	-
	THYR01001634	9.8 6.56	9. 15	14. 08 6. 39	5. 65	9.76	4. 62 2. 31	5.62 2.41	3.69 3.7	4. 86 6. 48		•		- 1
	THYR01001637	12.87	8, 05	12, 44	2.18 5.03	3.8 4.87	5. 49	3.32	2.86	2. 92		**		- 1
45	THYR01001641 THYR01001656	20.03	12.66	30. 89	7	10.07	5. 22	6, 32	7.53	5. 64	١.	**	_	- 1
	THYRO1001658	7, 58	6. 85	10.14	4.65	4.77	3. 94	3.38	2.8	2.5		**	-	_ 1
	THYR01001661	150.1	98.3	106. 51		166.2	125.8	74	80.32	53.76	ľ		1	
	THYR01001671	10.97	8. 43	12. 21	5	5. 04	3. 97	5, 12	5. 12	3.84	**	**	۱-	-
	THYR01001672	12.9		11.75	5.85	5. 91	5. 08	3.69	2.74	2.72	**	**	-	-
	THYR01001673	31, 04		26.79	16.85	19.7	19.9	22.8	23, 43	23.51	**	*	۱-	- 1
50	THYR01001677	8, 63	6.86	9, 97	3, 55	4.9	4. 2	5.4	6.55	6.85			۱-	
	THYR01001683	39, 34		25. 87	16, 11	18.7	19, 17	8.36	10,76	9.09	ı			
	THYR01001700	12. 31	7.03	9. 15	3.61	4, 23	3.04	2.79	3.63	4. 8	*	*	-	-
	THYR01001702	23.7	21.62	17.4	10. 57	5.8	11.44	6.01	4. 83	2.94	*	**	l -	-
	THYR01001703	7. 22	4. 23	5.14	2. 63	3. 23	1.81	2.37	2.42	2.04		*	-	-
55	THYR01001706	8.39		8.09	3.72	3. 11	3.37	3,24	2.65	1.26	*	**	-	~
	THYR01001721	11.69		16.73	6.09	4, 96	4. 46	3. 92	2.33	3.87	*	**	-	-
	THYR01001725	147. 69	144.4	144.74	111.99	128. 3	86. 63	130.43	106.61	93.76	*	*	-	-

THYRONOD1730   7.74   3.57   7.41   4.48   4.19   3.07   3.64   4.09   4.72   1.77   1.77   1.75						1	Table 5	03							
THYRRIOD1738   1,57 3,03 2,24   0,77 0,17 0,55 0,07 0,99 1,64   1   1   1   1   1   1   1   1   1		\$THYR01001730	7.74	3, 57	7, 411	4.48	4, 19	3.07	3.64	4.09	4.72		1		1
THYRONODIYAS  5,12 3,48 3,39 1,67 2,12 2,24 1,188 2,6 2,48				3.03	2.24	0.97	0.17	0.58	0.7		1.64		- 1	-	-
THYPRIOLOGYA6  0.25   1.84   2.0   6   1.81   4.55   9.12   1.27   8.09   11.05   8			5, 12	3.48	3.39	1.67	2. 12	2.94	1.98	2.6	2.48		- 1		ı
THYRRIDOLOTY70   20.01   12.02   16.57   10.05   13   10.49   5.77   6.07   5.43   1   -	5			14.84	20. 6		4.65	9. 25	12.7				- 1	-	-
THYROLOGITZ   20.01   12.02   16.57   10.05   13   10.49   5.17   6.07   5.43   1   -     -		THYR01001746	10.78	5.89	7: 69	2.83	3. 1	3.64				*		-	-
THYROID 01778   S. 03   S. 04   S. 04   S. 05   S. 0			20.01	12.02	16.57	10.05	13	10.49	5. 37						
THYRDIOLOTISS   9.15   5.26   5.98   4.02   4.49   3.27   4.45   4.57   5.77		THYRO1001772	29. 2	56.08	29, 11	28.47	34.34	15.95	9. 21						-
THYRDIO 1978   S. 68   S. 74		THYR01001778	39.03										*		-1
THYROUODIST    15.5   11.42   12.45   13.57   15.25   15.44   13.77   15.25   15.45   13.77   15.25   15.45   13.77   15.25   15.45   13.77   15.25   15.45   13.77   15.25   15.45   13.77   15.25   15.45   13.77   15.25   15.45   13.77   15.25   15.45   15.75				3.26	5. 98	4. 02	4.49	3. 37	4. 45	4. 26			- 1		1
THYRRIDO 1800   25.21   17.42   21.74   18.25   16.43   13.71   8.41   10.02   11.59   84	10	THYR01001796	66.96	62.34	52.36	31.61	31.58	30.06	74.83		78. 59	**	- 1	-	-
THYPRIOLOGIST   10.44   7.96   6.8   4.59   3.72   3.71   3.62   1.76   1.34   3   4   -			25.21	17.42	21.74	19. 92	16.43								
THYROTOLORIST   18.78   12.28   15.51   15.03   13.53   6.39   8.79   4.88   8.97   4   -		THYR01001803	12.96	11.11	10.61	7.04	8. 3	3. 44	5.71	5. 48				-	
THYPRIDIONISIS   14.15		THYR01001809	10.44	7.96	6.8	4. 59	3.72	3.71				*		-	
THYRIDIOLISIS   14.15		THYR0:001817	18,78	13.28	15. 51	15, 03	13, 53	6. 93							
THYROTOLOGIS   1.4   8.22   6.89   7.42   7.59   7.29   7.29   7.20   7.21   7.20	16			14.31	9. 69		6, 63	6.06	6.27			*	*	-	-
THYROHOUSES   50, 53   22, 88   37, 71   31, 71   34, 18   22, 81   22, 16   27, 13   21, 15   5   7   7   7   7   7   7   7   7	15		8.14	8.22	6.65	5. 42	7.94	4. 29	7.29	4, 26			- 1		-
THYROIODISS    8,86   7,83   3,92   4,14   4,55   2,55   3,7   10,44   3,46   THYROIODISS    7,70   31,41   1,24   9,48   10,56   7,02   4,02   4,01   4,05   4,0			50, 53	32.88	37.71	33.17	34. 18						*		-
TRACHIODODOS   9. 54   7.16   8.05   5.68   4.66   5.55   5.46   4.22   5.76   2 -   TRACHIODOD13   10.02   5.86   5.22   5.56   5.46   5.31   1.07   2.02   4.14   3   4 -   TRACHIODOD14   13.69   3.02   5.86   5.85   4.66   5.75   4.06   6.37   1.07   3.63   2   4 -   TRACHIODOD16   12.8   6.48   10.55   5.54   3.9   3.41   3.79   4.55   3.68   2.8   4 -   TRACHIODOD17   12.8   6.48   10.55   5.54   3.9   3.41   3.79   4.55   3.68   5.8   4 -   TRACHIODOL12   17.86   13.19   13.76   8.07   3.6   5.55   4.56   5.35   5.68   6.59   4 -   TRACHIODOL12   17.86   13.19   13.76   8.07   3.6   5.55   4.50   5.35   5.68   6.59   4 -   TRACHIODOL12   17.86   13.19   13.76   8.07   3.6   5.55   4.50   5.30   5.08   6.50   4 -   TRACHIODOL12   17.86   13.19   13.76   8.07   3.6   5.55   4.50   5.32   4.22   5.77   4   4   4   4   4   4   4   TRACHIODOL12   13.56   13.9   13.76   8.07   3.6   5.55   4.50   5.30   5.08   6.50   4   4   4   4   4   4   4   4   4   TRACHIODOL13   13.51   13.50   13.76   8.07   3.6   5.55   4.50   5.40   5.20   4.22   5.77   4   4   4   4   4   4   4   4   4			9, 86	7.83	3.92	4.41	4, 53	2. 55	3.7	10.54					1
TRACHICODOUS   1.05   5.28   5.2   2.58   4.45   5.1   5.07   2.02   4.14   5   5   -		THYR01001907	27.03	13.41	12. 49	9, 48	10.54	7.02							1
TRACHICODOTA   1.5 9   5.22   8.53   4.66   5.76   4.06   5.37   3.07   3.63   2     TRACHICODOTA   1.28   8.48   0.55   5.54   3.9   3.44   3.79   4.55   3.68   4.84   -   TRACHICODOTA   1.28   6.48   0.55   5.54   4.82   5.29   2.88   3.9   2.88   3.9   5.08   5.08   4.08   -   TRACHICODOTA   1.75   1.38   5.07   5.6   5.5   2   2.25   2.24   5.20   2.8   5.08   4.8   -   TRACHICODOTA   7.55   6.3   5.07   5.6   5.5   5.44   5.9   5.25   5.44   5.9   5.8   5.25   5.44   5.9   5.8   5.25   5.44   5.9   5.25   5.45		TRACH1 000006	9. 54	7. 16	8.08	5.68		5.53						-	1
TRACHIO00074   13,59   3,22   8,53   4,56   5,76   4,06   6,37   3,07   3,63   2   4   -   TRACHIO00072   13,85   2,22   2,10   15,55   5,54   3,9   3,41   3,73   4,55   3,66   2   4   -   TRACHIO00102   26,56   26,2   20,1   15,56   18,57   15,2   32,52   24,6   27,05   4   -   TRACHIO0012   17,56   13,9   3,16   2,55   3,46   3,36   5,08   6,8   4   -   TRACHIO0012   17,56   13,9   3,17   3,55   3,56   3,56   4   5   4   TRACHIO0012   17,56   13,9   3,17   4,55   3,56   5,56   4   5   4   TRACHIO0012   13,56   13,51   13,51   4,58   5,07   3,6   5,5   4,56   5,2   4,27   4   4   -   TRACHIO0014   31,51   45,86   50,29   14,82   25,81   22,15   25,29   16,85   2,76   4   4   -   TRACHIO0016   1,55   4,98   5,98   6,22   5,55   5,44   6,1   4   4   TRACHIO0007   32,64   30,11   26,81   15,7   21,99   18,56   37,52   29,77   33,83   4   -   TRESHIO00007   31,16   1,68   1,50   7,19   3,9   4,25   7,17   3,10   3,20   TRESHIO00007   32,64   30,11   26,81   15,7   21,99   18,56   37,52   29,77   33,83   4   -   TRESHIO00007   32,64   30,11   26,81   15,7   21,99   18,56   37,52   29,77   33,83   4   -   TRESHIO0010   4,76   3,22   5,71   3,93   4,25   7,17   3,10   3,90   TRESHIO0010   4,76   3,22   5,71   3,50   3,20	20	TRACH1000013	9.02	5.8	8. 2	2. 58	4.45	3.1	3.07					-	
TRACHIODOUZ   26.58   28.3   20.1   15.81   18.57   15.2   25.22   24.6   27.09   1     TRACHIODOUZ   17.36   13.19   13.78   8.07   3.6   6.8   5.91   3.9   3.9   2.8   5.9   5.08   6.9   1     TRACHIODOUZ   17.36   13.19   13.78   8.07   3.6   6.8   5.91   3.49   3.25   4.97   2     TRACHIODOUG   17.56   13.8   5.77   5.2   3.75   3.17   3.49   3.25   4.97   2     TRACHIODOUG   13.76   13.76   13.8   13.8   14.8   5.9   14.82   23.8   12.15   22.8   16.8   22.8   2.8   2     TRACHIODOUG   13.5   3.6   3.6   1.8   25.8   12.15   22.8   15.8   22.8   1.8   22.8   2.8   2.8   TRACHIODOUG   13.5   3.8   3.8   1.8   3.8   3.8   1.8   2.8   2.8   2.8   2.8   1.8   2.8   2.8   2.8   2.8   2.8   TRACHIODOUG   13.5   3.0   3.0   1.8   2.8   2.8   2.8   2.8   1.8   2.8		TRACH1000074	13.69	9.32	8, 63	4.66	5.76								
TRACHIODOID 8, 84, 85   85, 81   40, 40   22, 57   28, 39   20, 38   81, 35   50, 68   85, 91   TRACHIODOID 6, 77, 55   63, 85, 77   87, 87, 77   87, 87, 87, 87, 87, 87, 87, 87, 87, 87,		TRACH1000095	11.28		10.55	5. 54	3.9						\$\$		-1
TRACHIODOILS   17.58   13.19   13.78   8.07   1.98   45   5.91   1.48   3.25   4.79   2   4.70   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1.48   1.28   1		TRACH1000102	26.86	26. 3	20. 1							*	1	-	-
TRACHIGO0146 7.25 6.33 E.07 5.29 3.75 5.17 3.46 3.25 4.97 2 = - TRACHIGO0161 1.76 11.24 7.7 4.05 5.95 4.95 6.22 5.28 12.21 5.75 3 = - TRACHIGO0161 1.76 11.24 7.7 4.05 5.95 4.95 6.22 5.28 12.21 5.27 3 = - TRACHIGO0161 1.35 6.9 3.11 5.46 9.25 5.9 1.95 6.22 5.28 12.21 5.25 3.44 6.11 = - TRACHIGO0161 1.35 6.9 3.11 5.46 9.11 5.47 9.9 18.54 37.52 29.73 33.89 = - TRACHIGO0161 1.35 6.9 3.11 5.46 9.9 1.15 7.21 9.9 18.54 37.52 29.73 33.89 = - TRACHIGO0161 1.36 11.68 11.68 11.67 21.99 18.54 37.52 29.73 33.89 = - TRACHIGO0161 1.36 11.68 11.69 7.71 9.9 18.54 37.52 29.73 33.89 = - TRACHIGO0161 1.36 11.69 11.69 7.71 9.9 18.54 37.52 29.73 33.89 = - TRACHIGO0161 1.36 11.69 11.69 7.71 9.9 18.54 37.52 29.73 33.89 = - TRACHIGO0161 1.36 11.69 11.69 7.71 3.10 11.71 3.10 11.69 11.		TRACH1000108	88.56		40.48										-
TRACHIODOISIGN 11.78 11.24 7.77 4.05 5.59 4.89 5.32 4.22 5.87 3 2   TRACHIODOISIGN 3.51 54.86 5.05 91 4.82 25.81 22.15 25.28 16.88 22.15 2 -   VESSHIODOO04 11.55 6.9 8.13 5.46 9.50 5.81 22.15 25.28 16.88 22.15 2 -   VESSHIODOO031 3.16 11.68 12.05 7.19 3.9 4.25 7.18 4.46 8.92 3 4   VESSHIODOO051 3.17 20.53 22.45 10.12 5.91 10.79 18.54 3 7.72 2 2.77 3 3.33 1.96   VESSHIODOO059 4.76 3.32 5.577 3.56 3.32 3.56 1.16 3.33 1.96   VESSHIODOO107 158.28 125.83 134.47 126.46 135.27 13.26 1.16 3.33 1.96   VESSHIODOO107 158.28 125.83 134.47 126.46 135.37 133.6 74.81 79.38 82.68 4 5 4   VESSHIODO107 158.28 125.83 134.47 126.46 135.37 133.6 74.81 79.38 82.68 5 5 3 4.86 2 5 -   VESSHIODO107 158.28 125.83 134.47 126.46 135.37 133.6 74.81 79.38 82.68 5 5 3 4.86 2 5 -   VESSHIODO107 158.28 125.83 134.47 126.46 135.37 133.6 74.81 79.38 82.68 5 5 3 4.86 2 5 -   VESSHIODO107 158.28 125.83 134.47 126.46 135.37 133.6 74.81 79.38 82.68 5 5 3 4.86 2 5 -   VESSHIODO102 2 5.44 5.31 7.77 3 3 1.97 7.47 2.92 1.25 1.47 2.14 2.25 5 -   VESSHIODO102 3 1.44 1.02 1.02 1.02 1.02 1.02 1.02 1.02 1.02		TRACH1000126										*		-	
TRACHICODOM   1, 15   5, 16   5, 17	25	TRACH1000146													
VESENIODOCO		TRACH1000160												-	
VESENIODOOR   32.64   0.11   26.81   16.7   21.99   18.54   37.52   29.73   33.88   24.55												*		-	
VESENIODOS13   3.16   11.68   12.05   7.19   3.9   4.25   7.18   4.46   8.92   \$													*		- 1
VESSH1000029   23, 17   20, 53   22, 94   10, 16   15, 29   14, 76   9, 13   12, 72   8, 48   2   45   -     VESSH1000010   4, 76   3, 12   5, 75   3, 56   3, 22   3, 56   1, 16   3, 33   1, 16   5, 18   4   -     VESSH1000101   2, 76   3, 26   3, 27   3, 56   3, 28															-
VESENIODOSS    4.78   3.12   5.77   3.56   3.12   3.55   1.16   3.13   3.18															
VESENIODOSTO   8, 85   5, 54   8, 92   5, 55   8, 82   5, 44   8, 92   5, 55   8, 82   5, 44   8, 92   8, 95   8, 95   8, 95   8, 98   8   9   9	30											*	**	-	- 1
WESHI000171   St. 28   125.83   134.47   126.45   118.27   123.65   74.86   79.36   82.66   32.67   126.57															- 1
VESSH1000127   3,74															
*** **********************************			158.28										**		- 1
VESENIODOST  7, 93 10, 91 7, 7  8, 31 5, 46 4, 02 3, 34 1, 35 5, 77   2   4   -     VESENIODOST  2, 47 4, 75 1, 07 4, 43 4, 22 7, 07 3, 11 4, 32 11, 25 5   -     VESENIODOST  8, 95 56 93, 15 78, 83 74, 46 93, 71 81, 05 50, 08 36, 54 32, 24   24     VESENIODOST  8, 45 1, 46 1, 12 1, 19 3, 7 7, 86 1, 92 3, 92 6, 84 13, 85 21, 25     VESENIODOSS  40, 43 29, 46 56, 48 41, 45 2, 72 50, 13 17, 51 33, 82 20, 25     VESENIODOSS  4, 51 6, 13 9, 90 5, 55 84, 52 21 3, 15 57, 55 86, 64     VESENIODOSS  4, 51 6, 13 9, 13 1, 73 7, 84 56 5, 72 2 5, 82 6, 16 6, 5 2 2															-
VESENIODOSTS   12.47															
VESSH1000219   89, 55   89, 15   78, 83   74, 46   93, 71   81, 06   90, 08   36, 94   2,24   94   94   95   95   95   95   95   9	35												*		-1
VESHI000379   16.42   11.12   19.97   7.86   19.23   8.22   8.24   13.18   21.33       VESHI000380   4.5   6.13   9.09   5.56   8.45   6.21   3.44   6.29   2.57     VESHI000340   7.34   11.73   12.32   13.03   10.73   17.33   11.52   8.15   7.55   8.65   6.04   ** - VESHI000410   7.34   11.73   12.31   7.37   8.55   7.32   5.22   5.22   6.18   5.5   * - VESHI000410   4.34   16.7   17.37   8.55   7.32   5.25   6.4   8.25   7.07   ** - VESHI000411   4.34   16.7   17.37   8.15   6.25   6.4   8.25   7.07   ** - VESHI000411   4.34   16.7   17.37   4.25   4.15   5.15   5.25   6.4   4.28   7.07   ** - VESHI000415   6.55   6.57   6.57   6.55															-
VESENIOODSS  40, 42   23, 46   561, 48   41, 4   53, 72   50, 13   7, 75   31, 82   20, 25													**		1
## VESHI000388 4.51 6.13 9.09 5.56 8.45 6.21 3.44 6.28 2.57   ## VESHI000341 7.33 123.92 13.03 107.32 177.33 115.58 13.57 7.55 8.66 6.44 ** -   ## VESHI000410 7.44 11.73 12.31 7.37 8.85 7.32 5.82 6.16 5.5 9 -   ## VESHI000410 1.43 16.7 11.73 12.31 7.37 8.85 7.32 5.82 6.16 5.5 9 -   ## VESHI000410 7.44 11.73 12.31 7.37 8.85 7.32 5.82 6.16 5.5 9 -   ## VESHI000410 7.44 11.73 12.31 7.37 8.85 7.32 5.82 6.16 5.5 9 -   ## VESHI000410 7.01 14.64 7.07 14.00 1.00 1.00 1.00 1.00 1.00 1.00 1.0													i		- 1
VESENIO00394   17, 33   12, 32   13, 63   10, 12   17, 73   11, 55   8, 16, 57   78, 58   86, 04   ##   -     VESENIO0411   14, 34   16, 7   11, 87   14, 26   14, 7   9, 52   6, 54   4, 28   7, 07   ##   -     VESENIO0440   7, 1   4, 56   7, 48   10, 08   4, 59   9, 04   3, 62   6, 78   78, 77   ##   -     VESENIO0540   7, 1   4, 56   7, 48   10, 08   4, 59   9, 04   3, 62   6, 77   6, 78   78,															- 1
VESEN1000410   7,84   11.73   12.31   7.37   8.85   7.22   5.02   6.16   5.5   2   -	40														_ 1
VESHI000411   4.34   18.7   1.87   14.28   14.7   8.52   8.54   4.28   7.07   ***   -     VESHI000415   6.55   8.22   8.5   3.16   5.01   4.55   2.06   3.93   7.07   ***   -     VESHI000440   7.1   4.56   7.48   10.06   4.59   9.04   3.62   8.77   8.76   8.27   8.76     VESHI000539   87.4   47.68   87.32   38.3   45.7   42.77   51.56   51.15   50.18     VESHI000557   7.29   12.24   1.13   6.05   4.57   8.75   8.75   3.4   3.35   \$	40														-1
VESH1000415   6.65   6.22   8.5   3.16   5.01   4.35   2.06   3.39   7.37   1   5   7.74   1.06   6.45   5.90   4.35   2.06   3.39   7.37   1   5   7.74   1.06   6.45   5.90   4.35   2.06   3.39   7.37   1   5   7.74   7.68   7.68   7.74   7.68   7.68   7.74   7.68   7.68   7.74   7.68   7.68   7.74   7.68   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.68   7.74   7.75														1	
VESENIOOM401   7.1   4.56   7.48   10.06   4.89   5.04   3.62   5.77   5.76									0.04				**	_	-
VESENIODOS42  27.08 114.54   02.88   100.33   08.22   12.79   77.87   58.16   82.71   \$   \$   \$   \$   \$   \$   \$   \$   \$		AE2EM1000412										•			-
45   VESENIOOOSI9   S7.4   47.8   67.32   38.3   45.7   42.57   51.56   51.15   50.18				114 64											_ I
VESEN1000554   7.29   12.24   14.13   6.05   4.57   6.75   3.69   3.4   3.85   \$   \$   \$   \$   \$   \$   \$   \$   \$	45												•		
VESH1000557															- 1
VESENIOOOSTS   16   18.27   24.28   11.3   18.07   11.77   13.59   13.83   16.27													٠		
VESENIOODES  14.4   14.28   13.28   5.87   13.37   6.72   4.91   4.08   5.01   ±															- 1
*** VESENIOROSE2 28.09 20.04 22.75 8.98 11.53 77 7.53 8.48 5.44 28.29   **** VESENIOROSE3 9.27 5.38 6.39 4.39 2.39 3.11 3.3 12.18 6.06 #   *** VESENIOROSE3 7.65 11.52 16.09 8.2 9.11 7.91 4.21 6.96 3.61   *** VESENIORO752 22.77 18.91 24.8 11.19 11.63 9.8 14.24 19.48 18.19 22 -   *** VESENIOROSE3 7.03 8.3 9.3 5.64 5.4 3.65 3.8 14.24 19.48 18.19 22 -   *** VESENIOROSE3 17.03 8.3 9.3 5.64 5.4 3.65 3.1 8.25 5.7 2 -   *** VESENIOROSE3 16.55 20.4 16.66 7.12 12.23 7.45 5.22 4.66 8.91 2 22 -   *** VESENIOROSE3 16.56 20.4 16.66 7.12 12.23 7.45 5.22 4.66 8.91 2 22 -													**		٠.
90 YESH1000658 8.27 5.58 6.99 4.92 3.93 3.11 3.31 12.18 6.06 \$ - \ \text{VESH1000689} 7.65 11.52 18.09 8.2 9.11 7.91 4.21 6.96 3.81 - \ \text{VESH1000752} 22.77 18.91 24.8 19.46 23.17 19.77 12.01 15.14 12.53 \$ ** - \ \text{VESH1000757} 7.03 8.3 9.9 5.64 5.41 3.65 3.8 14.24 19.46 18.19 ** - \ \text{VESH2000039} 16.58 20.4 16.46 7.12 12.23 7.45 5.22 4.66 6.91 ** - \ \text{VESH2000039} 16.58 20.4 16.46 7.12 12.23 7.45 5.22 4.66 6.91 ** - \ \text{VESH2000039} 15.76 17.75 7.45 12.22 8.13 11.58 5.16 13.31 ** - \ \text{VESH2000019} 15.76 17.97 17.55 7.45 12.52 8.13 11.58 5.16 13.31 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.22 4.65 6.91 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 4.65 6.91 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 4.65 6.91 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 4.65 6.91 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 4.65 6.91 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 8.13 11.55 8.16 13.31 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 8.13 11.55 8.16 13.31 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 8.13 11.55 8.16 13.31 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 8.13 11.55 8.16 13.31 ** - \ \text{VESH2000019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 8.13 11.55 8.16 13.31 ** - \ \text{VESH200019} 16.59 20.4 18.66 7.12 12.23 7.45 5.25 8.13 11.55 8.16 13.31 ** - \text{VESH200019} 18.50 20.4 18.66 7.25 8.15 8.15 8.16 8.50 8.15 8.15 8.16 8.50 8.15 8.15 8.15 8.15 8.15 8.15 8.15 8.15												**		١.	
YESEH1000689   7.65   11.52   15.09   8.2   9.11   7.91   4.21   5.96   3.61   YESEH1000742   27.12   25.72   26.48   13.46   23.17   19.77   12.01   15.14   12.53   \$\$\$ = \$\$\$\$ - \$\$\$   YESEH1000752   23.77   18.91   24.8   11.19   11.63   3.6   14.24   19.48   18.19   \$\$\$\$\$   YESEH1000767   7.03   8.3   9.5   5.45   5.41   3.65   3.1   8.25   5.77   \$\$\$\$\$\$   YESEH2000039   16.55   20.4   16.46   7.12   12.23   7.45   5.22   4.66   6.91   \$\$\$\$\$\$\$   YESEH2000039   16.55   20.4   16.46   7.12   12.23   7.45   5.22   4.66   5.91   \$\$\$\$\$\$\$\$\$\$   YESEH2000103   15.76   17.97   7.75   7.45   12.22   8.13   11.55   5.16   13.31   \$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$	50												**		- 1
VESEN1000743   27.12   25.27   28.46   18.46   23.17   19.77   12.01   15.14   12.53   2 ± 2 - VESEN1000752   23.77   18.91   24.8   11.19   11.63   3.81   14.24   19.48   18.19   2 ± 3 + VESEN1000763   7.03   8.3   3.9   5.64   5.41   3.65   3.1   8.25   5.47   2 ± 3 + VESEN2000039   16.58   20.4   16.46   7.12   12.23   7.45   5.22   4.66   6.51   2 ± 3 - VESEN2000103   15.76   17.97   17.55   7.45   12.23   7.45   5.22   4.66   6.51   2 ± 3 - VESEN2000103   15.76   17.97   17.55   7.45   12.23   8.13   13.5   8.16   13.31   2 ± 3 - VESEN2000103   15.76   17.97   17.55   7.45   12.25   8.13   11.55   8.16   13.31   2 ± 3 - VESEN2000103   15.76   17.97   17.55   7.45   12.25   8.13   11.55   8.16   13.31   2 ± 3 - VESEN2001023   15.76   17.97   17.55   7.45   12.25   8.13   11.55   8.16   13.31   2 ± 3 - VESEN2001023   15.76   17.77   17.55   17.57   17.5												٠.		-	- 1
VESENIOOOTS2 22.77 18.91 24.8 11.19 11.53 9.8 18.124 18.19 18 - VESENIOOOTS1 7.03 8.3 9.3 5.64 5.4 3.65 3.1 8.25 5.77 8 - VESENIOOOTS1 16.55 20.4 16.66 7.12 12.23 7.45 5.22 4.66 8.91 88 8 - VESENIOOOTS1 16.78 17.75 17.45 12.23 7.45 5.22 4.66 8.91 88 8 - VESENIOOOTS1 16.78 17.75 17.45 12.22 8.13 11.5 5 8.16 13.31 8 8 - VESENIOOOTS1 15.76 17.57 17.55 7.45 12.22 8.13 11.5 5 8.16 13.31 8 8 - VESENIOOOTS1 17.55 7.45 12.22 8.13 11.5 5 8.16 13.31 8 8 - VESENIOOOTS1 17.55 7.45 12.22 8.13 11.5 5 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.5 5 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.5 5 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.55 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.55 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.55 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.55 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.45 12.22 8.13 11.55 8.16 13.31 8 8 - VESENIOOTS1 17.55 7.55 12.55 12.55 8.15 13.55 8.15 13.31 8 8 - VESENIOOTS1 17.55 7.55 7.55 12.55 8.15 13.55 8.15 13.31 8 8 - VESENIOOTS1 17.55 7.55 7.55 12.55 8.15 13.55 8.15 13.31 8 8 - VESENIOOTS1 17.55 7.55 7.55 7.55 7.55 7.55 8.15 8.15 8.15 8.15 8.15 8.15 8.15 8													**	١	. 1
VESENIO00761 7.03 8.3 9.9 5.64 5.41 3.65 3.1 8.25 5.47 * - VESENIO00029 16.38 20.4 16.46 7.12 12.23 7.45 5.23 4.66 6.91 \$ *** - VESENIO000129 15.76 17.97 17.55 7.45 12.52 8.13 11.35 8.16 13.31 \$ ***													**		- 1
VESEN2000039 16.58 20.4 16.46 7.12 12.23 7.45 5.23 4.66 6.91 * ** VESEN2000102 15.76 17.97 17.55 7.45 12.52 8.13 11.35 8.16 13.31 * *															-
55 VESENZO00102 15.76 17.97 17.55 7.45 12.52 8.13 11.35 8.16 13.31 * *													**	-	- 1
	55														-1
#ESCREUDU 104 03.00 07.00 07.01 10.02 (U. 0) 10.00 11.01 (U. 0) 4.01 10.00														۱.	- 1
		TESERZ 000 164	9 03.00	34.63	J4. 07	1 10.32	20. 39	(0.0	10.03	21.31	20.31		**	•	

				1		<i>J</i> 4 · ·							_
IVESEN2000175	7.45	6.95	7.39	3. 43	6.1	3.72	4.64	8. 1	4. 35	*		-	- 1
VESEN2000186	18.04	20.73	21.8	11. 19	14.77	10.58	8.74	12.84	12.61	**	**	-	- 1
VESEN2000199	11.71	6.48	9.96	8. 59	9, 24	5. 52	6.89	6.26	14.79		- 1		- 1
VESEN2000200 1	150.21		144.63			128.05	93.38	87. 48	97.05		**		-
VESEN2000204	7.05	7.07	9.65	4. 56	4. 24	3	2.19	4. 63	3. 19	*	*	-	- 1
VESEN2000218	16.65	12.07	17.29	7.31	8.24	6.16	8.32	5	5.15	**	**	-	- 1
VESEN2000230	12.09	13.48	13.57	4. 57	10.45	4, 08	3.93	4. 19	2. 98	*	**	-	- 1
VESEN2000272	8.95	9.01	11, 14	4.32	3.66	3. 22	4. 36	3. 53	4. 22	**	**	-	-1
VESEN2000299	8.61	7. 23	11.81	5.04	5.83	4, 25	3.6	5.71	4.76		* 1	-	- 1
VESEN2000323	17.3	12.74	15, 22	15. 36	17.47	14, 24	5.84	6, 35	4.65		**		-1
VESEN2000327	18.5	9, 19	14. 45	6.81	8, 17	7.45	3.2	5, 07	5. 85		*		- 1
VESEN2000328	7, 53	7.78	8. 38	3.45	4, 17	4, 12	2.35	2, 19	1.81	**	**	-	- 1
VESEN2000330	25. 07	20, 15	22.73	13.34	18, 45	13, 16	15.63	15, 15	15		**	-	-1
VESEN2000336	7. 29	9. 37	11.82	5, 86	9.09	5.78	3, 64	3.5	5				- 1
VESEN2000354	10.99	9. 78	10.4	5. 27	8, 22	4.93	3. 58	3. 58	2. 59	*	**	-	-1
			139, 59	81.44	99.69	74, 23	74. 17	76.59	54. 75	*	*	-	-1
					15.12	13	22.77	26, 45	21.84		`.I	_	- 1
VESEN2000379	32.08	21.47	28. 34	16.72		10, 8	15.72	16.01	17. 99		- 1		- 1
VESEN2000397	21. 17	10.11	25.4	12.77	12. 2	4.01	2.87	3, 94	3.48		*		-1
VESEN2000416	5. 92	5. 21	7.48	3.5	6.28			4. 27	1.92		٠,		- 1
VESEN2000420	8.47	4. 35	7. 18	4.04	3.81	2. 45	2. 31	2. 27	0.71		**		١,
VESEN2000430	8.74	5.76	7.33	2.94	5. 32	2, 11	1.8					-	-1
VESEN2000448	14, 67	14. 58	16.08	4.71	4.34	2.88	3. 91	2. 23	2. 27	**	**	-	-
VESEN2000449	46.1	50.29	39.97	32.75	32. 82	21.56	37.62	28. 99	23.73		. *	-	
VESEN2008456	45.88	33. 52	32.47	18.88	21.08	19.47	15. 48	22. 28	22, 31	*	*	-	-1
VESEN2000562	6.11	3.13	4. 69	5.4	3.95	3.19	3.24	2. 45	2. 42	١.			- 1
VESEN2000573	26.67	20, 52	18. 39	14.29	11.45	8.79	8. 94	7.46	12. 2	*	*	-	-1
VESEN2000604	9.02	7.76	4, 59	4.65	4. 32	4. 42	1.84	4, 41	1.34	1	*		-
VESEN2000614	9. 59	7.78	6.05	4.77	5.05	4.35	3	3.94	2.87		*	-	-
VESEN2000638	16.75	10.83	10.74	6.38	9.46	6. 26	4. 67	4.46	3.5		*		-
VESEN2000641	15.71	14.29	14, 25	7.95	7.6	6.71	8.94	7.41	6.6	**	**	-	- 1
VESEN2000645	9.28	11.51	10, 66	5, 27	8.01	4. 95	5, 42	4. 96	6.81		**	-	-
Y79AA1000013	24.66	15.88	15.09	6.08	5. 43	5, 53	13.59	13.07	18, 88			-	
Y79AA1000030	19. 52	16.62	14. 47	8.07	7.77	10.99	7.86	7.77	10.02	1	**	-	-
Y79AA1000033	29. 91	18.65	28. 38	16.37	15, 72	11.59	28, 15	19.64	28, 85			-	
Y79AA1000037	4. 94	2.8	4. 89	5, 26	3. 24	0.91	1.44	1.94	2. 43	i .		l	-
		53.74	54.67	39. 77	35.14	29. 21	42.77	30, 09	45.97	۱ *	1	-	
Y79AA1000041	79.98					2.06	5. 32	3,01	5. 11	1:		١.	-
Y79AA1000059	9.8	7.3	8. 44	4.37	5.08 124.96	111.44	68.74	63.93	65, 56	١.	**		-
Y79AA1000065	116.14		106.53	93. 49			5. 68	4. 38	3, 88	1	**		-
Y79AA1000081	7.88		9.72	7.03	9. 28	7.22		6. 34	4. 47		**	_	-
Y79AA1000127	8		6.2	3.17	4. 25	2.35	5. 28	16, 35	14.16			1-	- 1
Y79AA1000130			14. 68	13.04	8.43	8.05	14. 41						- 1
Y79AA1000131	17. 14		11. 18	6.93	10.63	6. 92	6.82	8. 55	6. 52		. !		
Y79AA1000134	6.34		5.68	7.86	5. 61	3.09	4. 16	2.2	3.07		*	ı	-
Y79AA1000143	8.89		8.14	3.93	5. 83	3.75	3. 23	2. 53	3.96		**	-	-
Y79AA1000144	11.79		9.18	5. 64	6.37	5.56	3. 97	3.84	3.36		**	١-	- 1
Y79AA1000150	28.57	28.05	20.04	12.4	17.77	10.86	24.45	16.09	20. 29			I -	
Y79AA1000153	11.32		7.41	5, 89	9.36	6.53	6, 84	5.59	6.52			ı	- 1
Y79AA1000166	13.75		9, 45	4, 16	5. 68	2, 88	7.42	8.85	8.7€			i	
Y79AA1000179			7.74		7.5	4, 67	13, 52	5.81	9.18			1	
Y79AA1000181	14.37		8.77	6.64	8, 41	3, 93	3.41	3, 97	4, 65			l	-
Y79AA1000202	22.9		19. 17		12.31	8, 36	18, 27	13, 59	20.94			١-	
			44.5		38.89	23.79	15.85	13, 31	14. 27		**	-	-
Y79AA1000207	52. 28 9. 44		5, 66		4.79	4. 18	5. 54	2, 28	3. 2				
Y79AA1000214					7.04	2, 26	4.7	3. 53	4, 67		*	1	_
Y79AA1000222			7. 95					16.69	21.48		•	١.	
Y79AA1000226					15.74		24.75					1 -	
Y79AA1000227	9.07		6. 62		4, 45		7.4	6. 45	7.51			1	
Y79AA1000230			29. 26		10.05	6.68	4.18	3.82	4. 93		**	1-	-
Y79AA1000231	65.27				47.89		28. 97	32.14	28, 29		*	1	-
Y79AA1000239					52.75		64.49	54.97	44. 91			1	
Y79AA1000258					4.01	3, 09	3.34	2. 93	4. 62	2		ı	
Y79AA1000268					30. 03		30.39	28, 87	31.5			1 -	-
Y79AA1000269								62.7	65, 45			1	
					7. 95				4.72			1	
Y79AA1000270					12. 9				6 20	5 **	**	1 -	-

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	Y79AA1000285	22.27	16.31	18.63	6.85	10.3	7.361	15, 47	22, 31	18.9	**	- 1	-	1	
										2. 22		- 1		- 1	
	Y79AA1000295	4, 35	2.68	5.01	1.59	3.76	2.11	2.3	2.59			- 1		- 1	
	Y79AA1000307	6.82	4.93	8. 121	6.39	9.3	7.36	3	5.88	4.49		ı			
5						3. 1	3. 25	1, 93	2.47	1.62		- 1		- 1	
	Y79AA1000313	4. 28	2.34	3.94	1.66							- 1			
	Y79AA1000314	1, 57	1.83	4.11	1.4	1.41	0.98	9.87	8.14	1. 73		- 1		- 1	
		3.78	3.39	5. 43	2.09	10.7	2.68	3.27	3, 33	3. 32		- 1		- 1	
	Y79AA1000328											1		- 1	
	Y79AA1000334	9. 59	9.06	7.49	5, 79	6.64	3.5	5. 12	3.62	4. 02	*	**	-	- 1	
	Y79AA1000342	17.48	13.29	18. 31	13.02	15.9	13.67	34. 2	31.79	30.43		**		+ 1	
												**		-1	
	Y79AA1000346	17.08	11.48	18. 69	8.72	9.82	6.74	5. 35	5. 97	4. 15	Ŧ		-		
10	Y79AA1000347	28, 82	26.71	31, 26	22.14	31.96	28.74	20. 9	20.82	19.69		**		- 1	
								5, 37	7.08	5, 15		**		- [	
	Y79AA1000349	15. 95	20.35	20.7	17.44	19. 4	18.58								
	Y79AA1800355	11.9	11.6	17.89	14, 98	13.32	9. 11	5. 25	6.54	7.95		*		-	
						6.47	3.77	2.77	3.21	1.95				- 1	
	Y79AA1000368	5.31	4.56	7.88	4.89										
	Y79AA1000388	9, 85	11.63	11.71	9.62	9, 75	4, 51	5. 83	4. 57	5.04		**		- 1	
	Y79AA1000392	13, 05	13, 68	11.25	7.92	12.06	6.34	12. 1	13.67	13. 52					
15												ì		- 1	
	Y79AA1000405	100.68	62.31	79.28	103.67	109, 59	81, 35	71.19	52.24	66. 44		1		- 1	
	Y79AA1000410	8, 13	6. 1	10.22	6.13	5. 36	5. 28	4.83	4.8	5, 17		- 1		- 1	
								6. 77	7.21	5, 64	±	**	_	-1	
	Y79AA1000420	11.1	11.77	13.46	5.41	8. 52	5.43							- 1	
	Y79AA1000423	7, 61	9.21	8.96	4.46	6.29	5.4	4, 36	2.75	3.4	*	**	-	- [	
		5, 75	6.96	10, 06	4,68	5. 08	3. 97	3.38	2.69	2.24		* !		- 1	
	Y79AA1000426											٠,		- 1	
20	Y79AA1000432	5.62	5. 18	10.2	4.71	5.75	4.03	2. 85	3, 99	8. 12		- 1		- 1	
	Y79AA1000453	32, 42	33. 52	37.01	20.21	25.21	18.7	36. 98	32.84	33.05	**	- 1	-	- 1	
										2, 11		- 1		- 1	
	Y79AA1000465	4, 43	1.91	3, 14	1.7	4. 29	1.76	2.57	2. 29			- 1			-
	Y79AA1800469	35. 31	28. 37	27.42	30.17	34, 28	32, 32	21.89	31,66	24. 15					
			8.51	13, 38	5. 22	5. 02	5, 9	5.84	10.93	8.28	*		-	ı	
	Y79AA1000480	11.78										1			
	Y79AA1000502	19.24	16.15	21.16	8.66	14	12.35	16.39	18.98	18. 17	*	- 1	-		
25	Y79AA1000521	75, 16	71.09	84.73	74.85	78. 28	67.85	48.04	51.73	45, 89		**		- 1	
20									11.71	11.75				1	
	Y79AA1000534	13.31	15. 15	20. 4	13. 25	14.05	10.71	13. 15				- 1			
	Y79AA1000538	12, 36	14.67	20.45	9.58	11, 27	5, 96	3.97	0	4.64		**		- [	
			16.32	14, 99	9.48	12.74	10.1	5.39	5. 2	6, 13		**	-	- 1	
	Y79AA1000539										•	***	_	ı	
	Y79AA1000540	10.2	7.44	12,08	3.41	€. 65	4. 35	12. 15	18. 44	10. 32		- 1	-		
	Y79AA1000560	180, 34	128.09	149, 28	162.08	194.06	150.66	118.75	125, 1	108, 26				- 1	
									16.21	10.56			-	- 1	
30	Y79AA1000574	12.15	9. 72	10.83	5.66	8.8	4.53	10. 59						- 1	
	Y79AA1000584	40.25	35.5	37.1	23.89	28, 68	25. 51	26. 96	34, 59	37, 41	**		-	- 1	
		9.65	4, 15	9, 45	2.79	3. 53	2, 1	3.54	8.38	4.6				- 1	
	Y79AA1000589										١	. 1		- 1	
	Y79AA1000598	34, 26	29, 42	37, 32	17.43	15. 95	16, 46	24. 99	27.55	21.09	**	*	-	- 1	
	Y79AA1000600		45.66	51.58	51.4	52, 23	39, 65	32.06	30.08	26, 05		**		- 1	
										7, 05	*		_	- 1	
	Y79AA1000609	7, 91	6.61	8.76	1.95	5. 66	3.3	3.74	4. 36				-	- 1	
35	Y79AA1000618	20.27	20, 13	21, 51	6,61	13.31	6. 56	19.86	14. 92	16, 11	**		-	- 1	
				18, 66	9.09	10, 58	9, 21	14.78	8.21	19,69	**		-		
	Y79AA1000627	18.35	16.53								**				
	Y79AA1000636	5, 21	2.52	7.03	3.36	4, 46	4.2	2.54	3.13	1.94					
	Y79AA1000649		21.35	25, 26	12.58	12.75	9.69	16, 99	19.63	19. 13	**		-		
										2.99	**		-	- 1	
	Y79AA1000656		5.14	7.46	2.86	2.97	2.28	2.83	3.72				-		
	Y79AA1800673	8.38	7.17	11.58	4.74	3.99	5.74	1.87	4.27	2. 18			-	- 1	
40	Y79AA1000674			127.73	143.89	140, 32	116.2	111, 11		120.08					
											١.		۱.	- 1	
	Y79AA1000678	22. 12	27.62	25.96	12.61	21, 55	13.08	21.7	19.94	19. 36				- 1	
	Y79AA1000682	14.62	11.86	13, 44	6.55	8, 13	6.38	12,74	13, 35	11.29	**		l -	- 1	
			24.6	33. 28	19.88	16.2	12.73	33.87	33.73	36. 17			1 _	- 1	
	Y79AA1000683												•	- 1	
	Y79AA1000697	46.01	35. 33	48.41	23.15	27.46	23. 15	15. 24	20.88	15.95	*	**	۳.	- 1	
	Y79AA1000700			5.68	3.34	3.06	2.62	3.53	2.27	2. 51	1		1		
45										15.36			1	-	
40	Y79AA1000702	50.6	33.01	36	23.71	24.96	18.05	18. 21	11.97		١.			~ (	
	Y79AA1000704	93.86	71.31	88. 5	118.92	93, 16	106, 68	74.22	61.46	46.95	1		1	. 1	
							2.09	2. 28	0. 26	2.39	**	**	I -	- 1	
	Y79AA1800705			7.37	1.82	2. 22					1 **		1		
	Y79AA1000717	8.36	6.77	8, 13	4.96	8, 96	4. 55	2.44	3.74	2.84	ı	**		- 1	
	Y79AA1000722			10.81	4.97	5, 79	3.98	6.28	5.41	5.45	**	**	۱-	-1	
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	Y79AA1800724	42.59	35.15	44. 52		30. 43	23.04	24.2	29.89	31.56	*		ı ~		
50	Y79AA1000726			7.31		4.77	4. 25	2, 41	3, 94	2. 09	ı	*	1	- 1	
								2.68	2.98	0.98			۱.	-1	
	Y79AA1000734					2.76	3.46						1		i
	Y79AA 1000748	7.45	7.72	8.69	2.89	2.82	2.75	2. 52	2.6	2.35	**	**	l -	~	i.
				33. 65		22.65	18.6		29.63	22.6	**		-		i
	Y79AA1800750							20.10					-	- 1	i
	Y79AA1000752	9.11	9.26	10.14	4.75	3.47	2.23		2.01	1. 29		**			i.
	Y79AA1000774		19.31	26, 89		13, 71	10.53	12.54	14.59	12.73			۱-	-	
55								1	10.24			-	١.	- 1	
	Y79AA100077	12.05	9.97	9, 69			2. 35			7. 92					
	Y79AA100077	17.61	11.59	13, 15	7.32	8, 15	7.21	6.02	5. 12	6. 88		*	١-	-	

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	[Y79AA1000778]	20, 06	13.62	11.66	10.8	9.59	8. 29	13.52	13.51	12.96		1		-1	
	Y79AA1000782	11, 43	7.55	9.19	7.76	8. 46	7.38	6.62	5. 21	4.82		*		- [	
5	Y79AA1000784	13.14	10. 21	12.84	4. 67	8.08	6.71	10.7	9. 8	9.71	**	- 1	-	- 1	
,	Y79AA1000794	46. 19	39.88	38. 82	22.35	24.71	16. 75	37. 28	33.24	34. 17	**		-	_	
	Y79AA1000800	7.18	3.91	5.09	0. 48 5. 78	0. 36 8. 8	0. 17 4. 96	2.34 3.89	0 4, 75	3.77	**	:	_	-	
	Y79AA1000802	11.45	13.12	10.85	5.78	11.14	4. 96	15.87	12.52	16.72	:		-	- 1	
	Y79AA1000805	27.49	21.74	19.65	16.59	17. 35	12.96	22	17. 22	22.17	•			1	
	Y79AA1000823	14. 24	7.73	9.85	4.94	4, 36	5. 06	6. 73	6, 96	8. 36	*	- 1	-	- 1	
10	Y79AA1000824	16.24	10	9. 76	8, 11	9, 28	6, 16	6.82	5, 83	8.72		- 1		- 1	
	Y79AA1000827	9.71	3.82	4.6	3.63	3, 22	2, 56	3. 51	2.06	2.83		- 1		- ŧ	
	Y79AA1000831	7. 38	5.79	7.27	4, 13	3. 32	2. 68	4. 43	0.57	1. 26	**	*	-	-1	
	Y79AA1000833	22.35	19.25	16.88	12. 38	7.9	8.89	19.38	11.86	18.04	**	- 1	-	ı	
	Y79AA1000850	33, 44	40. 57	26.99	19.52	21.81	14.62	28. 93	23. 5	32.01	*		-	- [	
15	Y79AA1000856	6. 25	7.4	4, 9	2.81	5. 29	2.12	4.31	2, 48	3.63		*		-1	
	Y79AA1000862	8.84	4.04	5. 05	7.04	4. 31	3.07	5.61	5.34	5. 05 6				- 1	
	Y79AA1000876	21. 17	10.25	11.73	7.67	6.82 17.29	6. 01 12. 59	5.79 18.41	7.01 20.33	23. 17		- 1	_	- 1	
	Y79AA1000888	32. 02 6, 26	25, 94 5, 68	21.68 6.17	17.64 4.27	4.4	2.72	2.73	2.97	2. 19	:	**	_	-1	
	Y79AA1000902 Y79AA1000935	76. 11	52.08	52. 32	70.65	71.58	69. 22	39.77	35. 2	32.3		*		- 1	
20	Y79AA1000959	48. 22	40.7	37. 08	25, 45	33.48	19, 11	26. 33	21.44	21, 93	*	**	-	-1	
20	Y79AA1000962	46. 82	45. 31	38.99	22.79	24, 29	17, 48	60.82	38, 12	53.14	**		-	- 1	
	Y79AA1000963	25. 33	35, 17	19.37	13.71	17, 32	12.79	14.75	14.39	13.92				- 1	
	Y79AA1000966	41.31	37.86	31.25	27.6	23,69	23. 7	32	18. 68	25. 22	*		-	- 1	-
	Y79AA1000967	19.13	9.69	10.03	8.88	7. 99	4.65	1.27	8. 2	0. 25				- 1	
	Y79AA1000968	25. 81	16.48	14.41	11.57	12.79	6.94	25. 2	19.6	28.68				- 1	
25	Y79AA1000969	40.02	39. 97	33.37	23. 55	39.74	36.04	32.93	30. 52	18.66				ı	
	Y79AA1000976	22.75	15. 17	18.99	14. 98	19.84	15. 88 2. 66	14. 85 4. 35	10.98	11.26		**	_	_ }	
	Y79AA1000978	8. 83 8. 02	7.09	6. 8 5. 83	4. 57	5. 39 4. 39	1.76	4. 37	2, 17	1.99	١.	••	_	-1	
	Y79AA1000985 Y79AA1000989			218.77	78.52	138.6	77.87	70.03	46, 49	61, 43	**	**	~	- 1	
	Y79AA1000991	12. 26	8. 35	10.64	5.63	5. 93	8.76	7.23	8.65	6.01				- 1	
30	Y79AA1001013	42.37	37.07	51. 31	24.72	27. 17	18.23	24.27	34. 59	22. 31			-	- 1	
	Y79AA1001014	16, 53	17.99	19.41	12.93	20. 24	14.74	13.35	18.92	15. 51				- 1	
	Y79AA1001019	8.07		8. 29	4.14	6.05	5.04	2.84	5. 3	3. 51		*	-	-1	
	Y79AA1001020	13.89	16.72	13.86	21.03	12, 53	10.76	9.5	10. 16	11.46		*		-1	
	Y79AA1001023	7. 28		9. 28	4.31	8.03	5. 29	3.01	3. 87	2. 51	•	:			
35	Y79AA1001030			33.86	36.62	21.08	22.74	16.86	22.85	18. 21 8. 91	1	:		-	
30	Y79AA1001035	10.6		10. 93 15. 3	9.68 8.73	10.73 9.98	7.69 7.28	7. 26	7.72 13.51	15. 31	١.	•	_	- 1	
	Y79AA1001041 Y79AA1001043	22, 79		22. 36	8.78	13.48	10.29	9, 42	13.86	17. 78	1 **		-	- 1	
	Y79AA1001048	12. 86		12.88	6.65	11.19	7. 59	6.44	7. 55	6.3		**	-	- 1	
	Y79AA1001056	6.67		12.37	5. 42	9.5	6.82	2.1	3. 99	2.72				- 1	
	Y79AA1001061	11, 12		15, 18	5.75	10.72	7.78	8.04	10.36	12.77	1 *		-	1	
40	Y79AA1001062	4.7		6.24	2.95	3.71	2.56	2.92	2.19	2. 28		**	-	-	
	Y79AA1001068			5.75	4.34	5. 98	4. 49	3. 29	3. 38	3. 07	١.				
	Y79AA1001073	17.6		20.08	5. 98	13. 24	7. 19	16, 17	13.71	20.42	*		-		
	Y79AA1001077	7.3		6.26	7.08	4. 43	3. 45	1.86	5. 22 5. 06	1.86				- 1	
	Y79AA1001078 Y79AA1001081	7. 29		6.25 8.85	3.73	3.81	2. 99 3. 52	2.8	3.1	3.74	1:		1	-	
45	Y79AA 1001088				24.5	22. 27	19	24, 36	30, 94	32, 13	**		۱-	-	
	Y79AA1001089					390. 54	406, 65	237.36	376, 46	392	1		1	ı	
	Y79AA1001090			8, 66	4, 12	3. 36	2.71	3, 39	1.8	2. 5	1		ı	-	
	Y79AA1001105				8, 51	14.07	9. 94	17.11	15. 32	15.62	**	**	-	-	
	Y79AA1001142				102.55	114.58	143. 24	69,56	60.81	74. 51	1	*	1	-	
50	Y79AA1001145	98.33		100.63		121.38	91.71	78	59. 2	74.76		**	l	-	
<i>S</i>	Y79AA1001162				92.38	93.62	73.32	97.56	124. 59	105.84			1		
	Y79AA1001167				7. 92	8. 62	6.56	7.15	16.96	9, 58		**		-	
	Y79AA1001176				76.52	65. 54	64.34	46.71	59.03 4.68	2.96		**	١.	-	
	Y79AA1001177				4. 06 28. 64	5. 51 33. 09	3.51 26.42	3. 45 13. 82	47. 33	10. 91		**	I -		
	Y79AA1001175					21, 34	16.32	19.9	17.65	25.06		**	-	-	
55	Y79AA1001201					31.35	17.81	48. 16	43.78	38. 95			-		
	Y79AA1001205					8.46	6.14		11.86	11.81			۱-	ш	1

Y79AA1001211	21 15	21.66	22.83	13.08	12.91	10.43	0.44	20. 18	19, 17	**		-	- 1	
										**	**		-	
Y79AA1001212			107.84			105. 91	80.87	83.57	67. 14					
Y79AA1001216	31.68	27.07	33.84	18.3	19.09	14, 63	17.05	18. 24	23, 12	**		-	- 1	
Y79AA1001228	7.97	5.25	12. 95	3.35	3.4	3. 64	3. 41	10.17	4.89		1		1	
									3, 13					
Y79AA1001233	8.26	6. 53	16.79		3.77	2.51	2. 28	3. 79			- 1		- 1	
Y79AA1001236	8.84	5. 25	11.14	5.02	5.53	3. 13	4.08	4.7	2. 24					
Y79AA1001239	74.22	59, 48	53. 54	45.62	83.11	43.05	50.35	42.44	46.18		- 1		- 1	
											1		- 1	
Y79AA1001240	70.83	49.07	64.4	30.83	35.04	20. 32	48. 49	59.48	54. 67	*		-		
Y79AA1001255	8.89	10.29	10.55	7.47	6,78	6.421	0.72	6. 35	0.76	**	*	-	- 1	
Y79AA1001264	14. 2	10.19	13.36	9.71	10.05	8, 67	4, 58	4.84	6.39		**		- 1	
									6. 53	**	**		-	
Y79AA1001272	15. 16	12.6	13. 5	3.77	6.76	5. 59	8.86	7.59				-		
Y79AA1001281	8.87	7.72	6.68	4, 17	4.8	3. 47	2, 65	2. 22	2.28	**	**	-	- 1	
Y79AA1001299	9, 39	10.08	9.6	4. 21	5. 3	4.49	3. 28	3.68	3.29	**	**	-	- 1	
											-	_	1	
Y79AA1001312	40.47	42.72	35. 81	17.92	27. 92	14.77	29. 7	20. 85	26, 67		7	_	- 1	
Y79AA1001319	31.92	23	20. 23	23. 19	28. 4	16. 96	30.43	20.02	181				- 1	
Y79AA1001323	32.99	27. 82	34. 32	14.07	19.03	14. 31	25.9	23.32	27.9	**		-	- 1	
										**			- 1	
Y79AA1001328	28. 33	16.1	28. 16	14.56	12.93	12.69	4. 94	7.51	4.42		*		- 1	
Y79AA1001343	12.56	8.01	10.99	6.43	7.59	7. 49	6.83	9, 08	8. 27					
Y79AA1001351	6.64	4, 21	8.23	2.99	1.8	1.8	2.08	2. 52	1,68			_	- 1	
										i		_	-	
Y79AA 1001364	8.06	5. 02	6.85	3. 26	2.98	2.12	4.06	2.93	1.74	•		-		
Y79AA1001367	145, 27	118, 42	143.45	117.7	132.8	94. 77	76.65	57.61	86.99		**		-1	
Y79AA1001384		52.17	41.89	27.43	34.83	21. 42	31.33	29.86	26.76	*	**	-	- 1	i i
											**		-	
Y79AA1001391	10.96	12.22	10.34	2.99	9. 55	4. 77	7.43	7.31	6.01		**		- 1	١.
Y79AA1001394	24.73	14.05	14. 28	9.7	6.32	9.07	6.42	10.16	9.01		- 1		- 1	
Y79AA1001402	12, 42	7.77	9. 12	5.62	4.8	4.67	4.74	3.87	4. 52	*		-	-	
		2.77	3.09	3.28		1. 95		0.78	1. 99		- 1		- 1	Ĺ
Y79AA1001410	5.24				1.53		2.62							
Y79AA1001414	27.49	18. 53	20. 44	10.78	9.47	10.2	16.16	13.43	11.92	*	*	-	-	1
Y79AA1001426		221, 23	225, 78	200.19	221.07	217.77	202. 63	211.55	197.64					
		10.97	10.9	6,77	10.81	5. 39	6.71	6.07	6.74				-	ł
Y79AA1001427	15.18													
Y79AA1001430	22. 25	17. 58	19.52	10.25	13.63	7, 85	11, 32	11.76	10.84	*	**	-	-	
Y79AA1001439	6.75	4, 58	4. 31	2.6	4, 23	1.25	2. 94	2. 35	2. 98				- 1	
		9. 99	14. 92	13.26	10.56	8. 25	5. 32	10.37	14. 36				- 1	
Y79AA1001485	17. 57												- 1	
Y79AA1001493	20.47	12. 23	13. 25	9.02	11, 27	6. 64	13.37	13.83	16.06					1
Y79AA1001511	14.52	8.05	9, 32	8, 33	8, 58	5.82	5.2	6	5. 95				1	
		6,72			4, 81	3.84	4, 54	3, 21	2.59				-	
Y79AA1001523											•		- 1	
Y79AA1001530	11.44	7.57	8, 59	6.64	8.84	5. 29	6.2	7.76	5. 77			1		
Y79AA1001532	13, 55	14, 73	11.04	5.35	5. 16	4, 93	5.08	4.52	6. 49	**	**	-	-	1
Y79AA1081533	56.5	68. 51	50.9	29, 97	36, 82	26. 92	69.28	61.19	56. 58	*				
												-		
Y79AA1001541	8.12	8. 52	5.25	4.02	4. 89	2. 24	4, 69	2. 24	2. 18		*		-	
Y79AA1001548	67.38	47, 55	44, 48	30,61	26, 42	21, 92	46.07	51.22	51, 86	*		-	- 1	
Y79AA1001555	14.35	9. 75		8, 86	4. 86	4, 97	8, 19	7.68	10, 98				- 1	ı
													- 1	
Y79AA1001562	18. 44	13.73		8.99	10.6	7.19	10.2	9. 39	15.75	*		-	- 1	
Y79AA1001581	24. 34	18. 91	17.62	9.8	12.04	10.4	15.9	9. 96	16.33	*		l -		ı
Y79AA1001585	30.32	20, 83		20.22	25, 12	15, 45	0.69	15, 13	0.14				-	
									14.72			١.	-	
Y79AA1001592	28. 53	25. 65		29.33		7. 28	20.69	8. 44			٠		- 1	
Y79AA1001594	23.28	21.85	24. 28	14.25	15, 59	10, 11	22.55	14.53	20. 27	**		۱-		
Y79AA1001603	12, 47	11.04	10.84	6.8	11.48	6.78	22.63	14.33	27.5					
Y79AA1001613		18.27				13.34	20.3	16.35	18, 99	1		1		
										1		ĺ		
Y79AA1001630		13.91				8.02	11.73	12.98	17.67			1		
Y79AA1001647	33.54	24, 86	22.66	23, 22	25.84	16, 49	26.93	21.3	26. 55			ı		
Y79AA1001664		10.56		7.78		5. 54	10.34	8.1	11.69	**		-		
									7.69			١	-	
Y79AA1001665		9				6.13	5, 92			*	*	١-	- 1	
Y79AA1001679	24, 51	18, 98	17.88	11, 13	12.84	9, 27	24.02	13. 28	18.8	*		l -		
Y79AA1001592		6, 72		2.24		2, 53	3, 64		3.06	**	**	-	-	ı
							14. 27	13.28	13.01			١_		
Y79AA1001696						4.61				١.		1 -		ı
Y79AA1001705	12.04		8.55	9.76	11.8	6. 21	4, 61	5.98	4.57	ı	**	1	-	ı
Y79AA1001711	13,72		10.63	6, 49	4.11	3, 08	6.08	4,06	7.4			l -		
							1.49		1.39	1		1		
Y79AA1001717		1.55				0.7				ı		ı		
Y79AA1001719	22.12				12.72	6. 53	12.21	11.1	11.76	1				ŧ
Y79AA1001727						9, 31	38, 01	28, 39	34, 23			I -		
Y79AA1001750						8.01	11.94		11.41		**	۱-	_	ı
											**		- 1	ı
Y79AA1001765	127.24	115,03	77.78			43.62	140, 25					1 -		
Y79AA1001777	6.77	6. 32	6.37	2, 57	5, 24	2,46	5.26	4. 15	4.71		**	۱-	-	
Y79AA1001781						2.55			1. 3			1	_	

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	Y79AA1001787	5, 04	3, 62	6.16	2.83	4. 49	3. 23	3. 33	2.97	2.01		- 1		- 1	
	Y79AA1001793		97.84	90.95		108.02	70.38	56, 01	50.79	40, 18		**		-1	
	Y79AA1001795	5. 66	5.4	6, 21	4.2	5. 9	3.1	3.91	2.49	2.28		**		-1	
5	Y79AA1001799	15.09	14. 95	16.22	8.14	9, 02	8, 13	5. 96	8.67	5, 96	**	**	_	-1	
	Y79AA1001800	25. 64	16. 41	26.8	14.68	31.06	13.67	17.04	13.04	14.7				- 1	
								10.02	6. 35	8.81		. 1		- 1	
	Y79AA1001801	18.54	13.81	14. 73	8.69	14. 72	7.61					٠,		- 1	
	Y79AA1001803	4. 48	5.85	6.44	4. 5	2.73	1.98	4.84	2.96	2.69		1		- 1	
	Y79AA1001805	40.28	26.89	34.51	17.56	11.83	10.76	12.11	14.87	11.59	**	**	-	-1	
	Y79AA1001807	42.38	40.49	41.45	39.87	44. 56	37. 42	30.19	22.76	22. 12		**		- 1	
10	Y79AA1001827	9.32	11.28	9, 81	8.01	8, 55	6.68	5, 41	8.86	5.83	*	*	-	- 1	
	Y79AA1001846	22.14	19.33	18, 59	11.82	17.37	13.61	12.56	15.77	9, 7	*	*	-	- 1	
	Y79AA1001848	8.04	9. 99	9, 11	5.33	6.98	5. 01	6.65	6, 63	5, 92	*	*	-	- 1	
				9. 06	5.77	8. 29	4. 56	4.63	5. 75	4, 91		**	-	- 1	
	Y79AA1001853	10.87	10, 1							18.07	•	71		-1	
	Y79AA1001863	24.97	27.27	22.17	17.04	22	11.97	13.95	21.02					-	
15	Y79AA1001866	13.46	10.72	11.93	6. 28	4. 52	4. 93	6.56	4.65	4, 51	**	**	-	- 1	
10	Y79AA1001874	2.1	1.41	2.93	0.78	1.08	0. 95	1.46	1.22	1.08		- 1		- 1	
	Y79AA1001875	31.78	22.74	37.59	12, 11	19, 59	16,03	9.16	15	14, 61	*	¥.	-	-	
			778.32				821.68	479 34	605.65	529.63		- 1		- 1	
	Y79AA1001908	5. 27	6.69	9, 96	3, 58	5. 35	3. 25	2.93	3.09	1, 48				- 1	
		8. 48	16.75	14.8	9. 28	7.65	6. 34	6.48	6.75	5. 36				-	
	Y79AA1001923											**		-	
20	Y79AA1001927	18.05	21.6	19. 1	12.29	16. 25	12. 12	14. 45	14.01	14.79	٠		_		
	Y79AA1001930	12. 23	24. 48	15.86	10. 91	18. 16	6,72	4, 54	5. 11	5. 37		*		-1	
	Y79AA1001932	61.74	56.58	52.19	37.44	49.06	33.54	58. 52	51.29	51.59	*		-	ł	
	Y79AA1001933	9. 52	6.51	10, 88	6.04	6.05	6, 44	4. 27	11.51	4.91		1		- 1	
	Y79AA1001942	10.76	6.66	7.93	7.97	5, 94	4.4	3.29	2.7	3.99		*		- 1	
	Y79AA1001963	138. 12		117.87	95.52	130. 54	91.9	83.12	67.08	69.04				- 1	
			91.21	62.32	82. 32	49.63	76.26	20.14	2.41	10.4				-1	
25	Y79AA1001968	147.27					10.20		3.73	3, 02		**	_	- 1	
	Y79AA1001983	12.1	16.25	14. 99	5.06	8. 59	7.83	3.08	4. 84	4, 51	7	*		-	
	Y79AA1002000	8. 87	9.86	16.46	6. 28	9. 55	3.45	4.66						- 1	
	Y79AA1002004	46.17	58.66	37.42	34.61	47	15. 44	17, 13	16. 13	8.67		**		- 1	
	Y79AA1002008	16. 52	21.1	20.83	11, 85	16. 14	12.02	7.95	15, 74	18.89	*		•	- 1	
	Y79AA1002012	13.85	12.54	10.41	5, 85	6.84	6, 65	8.76	11.21	9, 52	**		~	- 1	
30	Y79AA1002017	15.21	10.94	15, 42	6.73	12.23	8. 22	1.29	15.64	2.06				- 1	
30	Y79AA1002022	42.57	32.77	37. 11	21.13	22.85	16. 91	21.09	21.59	19, 22	**	**	-	~ 1	
			8.04	15. 25	2. 11	4.6	4. 84	5. 16	2.67	4.2	*		-	- 1	
	Y79AA1002027	10.35							2.07	7. 33	1	**	1.	_	
	Y79AA1002050	13.21	11.11	14. 39	7. 26	8.9	5. 9	6.9	7.16		٠.	**	1	- 1	
	Y79AA1002058	167.29	130.21	156.91	193.04	186.39	149.35	127.37	93. 56	137.15				- 1	
	Y79AA1002060	54.35	76.53	61.68	36.08	41.21	29, 16	24.43	9.99	5.28	*	**	-	- 1	
35	Y79AA1002062	49, 46	32, 53	50.05	16.65	20, 72	17.6	35.11	28. 1	35.48	*		-	- 1	
	Y79AA1002065		69, 34	64, 62	65.01	102, 96	72.96	82.29	90.04	44.91				- 1	
	Y79AA1002067	33.46	37.19	43. 9	21.85	20.1	22. 68	15.4	10, 55	10.37	**	**	-	- 1	
	Y79AA1002069	5, 33	3.94	7. 12	3.04	1.49	2.47	3.1	3. 24				-	- 1	
		67.39	142.78	83.33	77.38	149, 87	86, 41	153.3	88, 92	78.17	1		ì	- 1	1
	Y79AA1002070								491, 49				ı	- 1	
	Y79AA1002074	1225	1102	498. 16	721.59	1162.7	771.32				١.			- 1	1
40	Y79AA1002076	8. 24	11.39	15.58	4.01	6. 13	3, 99		9. 01	11.16	*		١-	- 1	ı
	Y79AA1002083	11.94	6.94	9.48	4. 49	4. 53	3. 15	4.48	1. 37		*	*	-	-1	ı
	Y79AA1002084	20.05	13.84	17.88	7.41	8.36	6.24	10.42	8.8		**	*	-	-1	i
	Y79AA1002086	15. 57	8.01	9.58	4. 16	3.65	3.41	3.6	4. 3			*	۱-	-1	
	Y79AA1002087	265.36	258, 38	345.6	296.44	355.85	246. 9	347.62	322.78	317.34	1		•		ı
	Y79AA1002089	15.8	12.04	15.08	6.67	9, 17	6, 14	12. 29	9.32	9, 91	**		-		
45	Y79AA1002093	13.66	12.64	9. 33	5, 35	6.21	4. 47	3. 01	2.8	2.57	**	**	-	-1	ı
40			7.04	8. 08	3. 07	2.77	1.96					*	١.	-1	i
	Y79AA1002101	6.58										**	1:	-	ı
	Y79AA1002103	11.98	11.79	10.9	4.11	5. 96	4. 92					**	_	-	ı
	Y79AA1002115		18. 46	14. 58	9.3								-		ı
	Y79AA1002121	6.93	5. 28	6. 92	6.7		4.36					**	1	- 1	ı
	Y79AA1002125	40.85	21.04	26.11	14, 22	18.4	17.66	12.21	8. 56				1	-	ı
50	Y79AA1002129		7. 25		4, 36	3. 94			3.8	4.37	١.		l -	-	ı
	Y79AA1002131	5.89		4.65									۱ -	- 1	ı
												**	1_	_	ı
	Y79AA1002139												-		ı
	Y79AA1002144	53.23	47.33	47.13		42.65						**	-	-	ı
	Y79AA1002177	14.09	13.17	13.65	5, 87	6.08	5. 05	5. 82			**	**	- 1	-	ı
	Y79AA1002183		99, 47		24.87							**	1 -	-	ı
55													١-		ı
	Y79AA1002202												1		ı
	Y79AA1002204	4. 17	2.23	3. 23	5.04	1.05	2, 68	3. 28	2.	4 2.69	1		1	- 1	•

	<del></del>
Table 509	

						Lucio 5	0,								
	1Y79AA1002206	7.86	6.79	4.86	2, 41	3.6	5.74	5.49	1.51	2.8		- 1		- 1	
	Y79AA1002208	21.91	17.64	15, 14	5.6	4, 07	5. 57	6.81	5. 17	2.55	**	**	-	- [	
5	Y79AA1002209	14, 82	11, 28	11.86	6.23	4.7	2. 82	4.71	1.33	3, 18	**	**	-	- 1	
	Y79AA1002210	13.64	7. 39	7, 59	9.08	4, 62	5, 18	20.5	2.05	6. 37		- 1		1	
	Y79AA1002211	11.76	19.59	13:47	10. 43	6, 65	6, 52	12.32	8.42	11.25		- 1		- 1	
	Y79AA1002213	40.78	31.99	22.96	18, 41	26. 57	14. 98	45. 88	32.4	41.97		- 1		- 1	
	Y79AA1002215	54, 92	41.69	39. 55	24.88	24, 36	11. 26	37.49	23.6	35.98		- 1	-	- 1	
	Y79AA1002220	17.03	11.5	20.58	7. 13	5. 68	5, 31	4, 57	4.8	6. 51		*	-	-1	
10	Y79AA1002226	48. 55	31. 27	31.34	7. 35	12,72	13.65	9, 19	6.65	11, 13	*	**	-	- 1	
	Y79AA1002229	7.88	6. 84	5. 37	6.02	4.67	2.85	3.67	2.52	3.73		*		-1	
	Y79AA1002234	20.83	13. 27	12.39	9, 34	6. 36	3.6	6.9	3, 36	5.84	*	*	-	-1	
		28. 03	23.84	21. 24	15.07	14. 87	9. 39	10.75	8. 42	13.64		**	-	- 1	
	Y79AA1002235	9.72	14.9	10.35	5. 25	6.99	4. 12	8.31	3.82	6.69	*	- 1	-	- 1	
15	Y79AA1002246	12, 35	12.02	7. 88	7.82	11.57	8. 55	9.77	4.5	4, 81				- 1	
	Y79AA1002258		49. 19	41. 11	5.28	2.78	2. 07	15, 99	20.03	22. 13	**	**	_	-1	
	Y79AA1002279	51.52	7.58	4.14	5.73	4, 43	2.94	6.45	6.01	8.36				- 1	
	Y79AA1002292	13.64	8. 29	4. 77	3. 29	5, 48	4. 2	4. 42	4. 85	3.79		1		1	
	Y79AA1002298	9. 43	8. 07	6, 62	3. 2	4, 4	2.44	5, 17	1, 76	2, 58			_	- 1	
	Y79AA1002307	9. 31	7. 96	8.79	3. 32	4. 26	2.83	4. 23	2.88	3, 97	**	11	_	- 1	
20	Y79AA1002309	8.88	9. 27	11.76	6.9	6.2	3.45	7.45	3, 09	6.4	*		-	- 1	
	Y79AA1002311	16. 51	8.36	7.7	5. 13	4, 89	3. 36	5. 91	4, 32	5.92	•		-	- 1	
	Y79AA1002334	13.05	12, 49	9. 42	7.1	4, 15	6. 19	7. 95	4, 93	5. 8			-	- 1	
	Y79AA1002351		30.6	21.85	12, 21	15.81	9.74	20.54	18. 29	18, 48	*	*	_	- 1	-
	Y79AA1002355	31.74	15. 4	18, 02	12. 53	10.73	6.85	25.86	17.1	25. 5			-	- 1	
25	Y79AA1002361		6.37	7.19	3. 15	4. 11	3.03	4. 29	4.74	4, 01				- 1	
		8. 95	6.89	5. 46	5. 13	4. 81	3.4	9. 6	4. 57	7.84				- 1	
	Y79AA1002373		2569. 2		462, 62		615.71		1040.3	1062.1			-	- 1	
	Y79AA1002376 Y79AA1002378	20.24	17. 32	13, 54	5. 14	9.41	4, 23	19.28	11.07	16.88			-	-	
	Y79AA1002378		128.86			141.78	92.68	155.95	123.08	170, 94			i		i
	Y79AA1002388	33.4	33. 3	27. 31	13. 85	26.75	11.62	21.29	16.32	21.24	1			-	1
30	Y79AA1002399	11.13	8. 22	7.72	4. 28	5. 54	4. 87	7, 56	5. 9	6. 25			-	1	
	Y79AA1002407	12.66	14. 43	18. 13	7.72	14, 18	6.84	5. 83	9.78	4, 59		*	l	-	
	Y79AA1002407		12.77	14. 95	5. 14	9, 13	4, 62	8.44	10.73	7.99		*	-	-	
	Y79AA1002416		8. 19	8.76	5. 47	10.72	5. 8	8.2	6.05	6, 59			1	- 1	1
	Y79AA1002410		18.61	8.81	5. 82	10. 24	4.73	3.65	6, 89	5.66	1			-	ı
35	Y79AA1002431	3.38	3.05	6. 2	3.01	5. 89	1.6	2. 81	2.79	1.69	1		ì		ı
	Y79AA1002433			9, 29	5, 11	5. 57	3, 18	3, 49	4.6	3.87	**	**	-	-	
	Y79AA1002445		25.62	23.49	15.99	10.67	7.02	18. 92	25.26	13, 87			-		l
	Y79AA1002461			7.84	3, 36	7. 35	4.7	3, 49	2.25	3, 85		**	l	-	l
	Y79AA1002466		130 4	681.02		499, 15	369	592.67	971	768.71	1		1		
	Y79AA1002471			15.35	12.81	13.4	11, 43	4.94	6.06	4.47	1	*	1	-	1
40	Y79AA1002472			31.17	18.15	21.85	9.34	16.29	20.14	20.03	1 *	**	-	-	ı
	Y79AA1002474			10.71	6.77	7.3	7.75		7.37	4.86	**	*	1 -	-	
	Y79AA1002482			36.63	19.02	23, 45	17. 38		25, 81	23.08	**	*	-	-	ı
	Y79AA1002487			7.43	7.28				3.78	3.86		**	1	-	ı
	Y79AA1002490				56, 22		57. 31		51.39	52.37	**	**	1 -	-	ŀ
45	Y79AA1002493			40.36	20.64				46, 19	46.7			1 -		ı
	ZRV6C1006278			5. 52	3, 16		2. 19		2.6	2.72	**	**	1-	-	ı
			· · · · ·												-

### EXAMPLE 16

Selection of novel cDNA clones from cDNA libraries prepared by oligo-capping method

[0246] The following 54 clones were newly selected from cDNA libraries prepared by oligo-capping method, based on the criterion that the 5-end sequence of a cDNA done contained a coding region which was initiated with ATG codon and which encoded 50 amino acids or more:

HEMBA1000497, HEMBA1001750, HEMBA1003854, HEMBA1004193, HEMBA1004860, HEMBA1005572, HEMBA1006038, HEMBA1006092, HEMBA10060406, HEMBA1006600, HEMBA1006601, HEMBA1006601, HEMBA10060812, HEMBA10060812, HEMBA10060814, MAMMA1002094, NT2RM4000634, NT2RM4000657, MAMMA1002094, NT2RM4000634, NT2RM4000654, MAMMA1002094, NT2RM4000654, MAMMA1002094, NT2RM4000654, MAMMA1002094, NT2RM4000654, MAMMA1002094, MAMMA1002094

NT2RM4000783,	NT2RM4000857,	NT2RM4001178,	NT2RM4002420,	NT2RP2000198,	NT2RP2000551,
NT2RP2000660,					
NT2RP2001214,	NT2RP2001460,	NT2RP2001756,	NT2RP2002056,	NT2RP2002677,	NT2RP2002755,
NT2RP2002843,	NT2RP2003101,	NT2RP2003799,	NT2RP2004095,	NT2RP2004732,	NT2RP2004920,
NT2RP2005454,	NT2RP2005776,	NT2RP2005806,	NT2RP2005882,	NT2RP3001282,	NT2RP3001723,
NT2RP3002099,	NT2RP3003155,	NT2RP3004028,	OVARC1000008,	OVARC1000724,	OVARC1000751,
OVARC1001029,	PLACE1000814, PL	ACE1003030, PLA0	CE1005549, PLACE	1007218.	
Among them, the	following 23 clones	was predicted to co	ntain a coding region	n encoding 100 am	ino acids or more:
HEMBA1000497,	HEMBA1003854,	HEMBA1004193,	HEMBA1006812,	HEMBB1001871,	NT2RM4000657,
NT2RM4001178,	NT2RP2001756,	NT2RP2002677,	NT2RP2002755,	NT2RP2002843,	NT2RP2004095,
NT2RP2004920	NT2RP2005806	NT2RP3001282	NT2RP3002099	NT2RP3003155	OVARC1000724

[0247] Table 510 shows maximal ATOpri value determined for each clone. Since the respective maximal ATOpri values for HEMBA1008B12, HEMBB1001B71 and NT2RRP30012B2 are higher than 0.3, the clones would be fulllength. Other clones indicated below have maximal ATOpri values of 0.3 or less, and this means that the fullness ratios

OVARC1001029, PLACE1000814, PLACE1003030, PLACE1005549, PLACE1007218, This indicates that the clones

However, the sequences can still be full-length:HEMBA100497, HEMBA1001750, HEMBA1003854, HEMBA1004193, HEMBA1004860, HEMBA100572, HEMBA1006038, HEMBA1006092, HEMBA1006606, HEMBA1006605, HEMBA1006605, HEMBA1006605, HEMBA1006605, HEMBA1006605, HEMBA1006605, HEMBA1006605, HEMBA1006605, MTZRM4000783, MTZRM4000657, NTZRM4000178, NTZRM4000657, NTZRM4000657, NTZRM4000657, NTZRM4000657, NTZRM4000660, NTZRM2000660, NTZRM2001740, NTZRM2001796, NTZRM2001796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZRM2000796, NTZMM2000796, NTZMM2000796

[0248] Table 511 (same as Table 2) shows SEQ ID NOs of the nucleotide sequences located at the 5'-end and 3'end of each of the 54 clones and the corresponding plasmid clone, which was obtained herein, containing a polynucleotide as an insert. SEQ ID NO for a 5'-end sequence is indicated on the right side of the corresponding Sequence name of 5'-end sequence, and SEQ ID NO for a 3'-end sequence is indicated on the right side of the corresponding Sequence name of 3'-end sequence.

[0249] Swiss-Prot was searched for data homologous to the 5'-end sequences of the selected 54 clones, and Gen-Bank and UniGene were searched for data homologous to the 5'-end and 3'-end sequences of the same clones. The search results are indicated as Homology search results 1-7 in the last part of this SPECIFICATION.

- Se [250] Based on the matching data obtained by the search, 7 clones presumably encode proteins belonging to any of the categories of sererotory or membrane proteins, glyaporteins, signal transaction-associated proteins, displayed proteins, and protein synthesis- and/or protein transport-associated proteins, displayed by the monology to any of known proteins belonging to said categories. Here, the term "relatively low homology (which means that when the nucleotide sequence exhibits "relatively high nomology" (which means that, when the nucleotide sequence is compared with the known sequences in Swiss-Prot database, the sequence in the sequence in Swiss-Prot database, the sequence in the sequence in Swiss-Prot database, the sequence in the sequence in Swiss-Prot database, the sequence in the categories and that, when the nucleotide sequence is compared with the known sequences in Swiss-Prot database, the sequence in the categories and the sequence in the sequence in the categories and the sequence in the sequence in the categories and the sequence in t
- 45 [0251] Among the 7 clones, clones presumably encoding proteins belonging to the category of secretory or membrane proteins are the two clones. HEMBE 1001871 and NTZRMA000857 (which also belong to other categories); clones presumably encoding proteins belonging to the category of glycoproteins are the two clones, HEMBE 1001871 and NTZRMA000857 (which also belong to other category of glycoproteins are the two clones, HEMBE 1001871 and NTZRMA000857 (which also belong to other category of signal transduction-associated proteins is PLACE 1005549; clones presumably encoding proteins belonging to the category of transcription-associated proteins are the three clones, HEMBE 1005572, NTZRP2001756, and NTZRRA000857 (which also belong to other category of disease-associated proteins is NTZRMA000857 (which also belong to other categories); a clone presumably encoding a protein belonging to the category of protein synthesis and/or protein transport-associated proteins is HEMBE 1001756 (see Examples 12).

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encode proteins.

Table 510
The maximal ATGprl value of each clone selected in Example 16

clone name		maximal ATGpr1 score	
HEMBA1000497	F-HEMBA1000497	0. 25	
HEMBA1001750	F-HEMBA1001750	0.08	
HEMBA1003854	F-HEMBA1003854	0, 23	
HEMBA1004193	F-HEMBA1004193	0.22	
HEMBA1004860	F-HEMBA1004860	0.29	
HEMBA1005572	F-HEMBA1005572	0. 24	
HEMBA1006038	F-HEMBA1006038	0.29	
HEMBA1006092	F-HEMBA1006092	0, 28	
HEMBA1006406	F-HEMBA1006406	0. 26	
HEMBA1006650	F-HEMBA1006650	0. 22	
HEMBA1006812	F-HEMBA1006812	0.71	
HEMBB1000672	F-HEMBB1000672	2 0.24	
HEMBB1001197	F-HEMBB1001197	0.22	
HEMBB1001871	F-HEMBB1001871	0.94	
MAMMA1001252	F-MAMMA1001252	0. 29	
MAMMA1002094	F-MAMMA1002094	0.28	
NT2RM4000634	F-NT2RM4000634	0.07	
NT2RM4000657	F-NT2RM400065	7 0.24	
NT2RM4000783	F-NT2RM4000783	3 0.22	
NT2RM4000857	F-NT2RM400085	7 0.12	
NT2RM4001178	F-NT2RM4001178	0. 27	
NT2RM4002420	F-NT2RM400242		
NT2RP2000198	F-NT2RP2000198	3 0.15	
NT2RP2000551	F-NT2RP200055	1 0.07	
NT2RP2000660	F-NT2RP2000666	0. 22	
NT2RP2001214	F-NT2RP200121	4 0.26	
NT2RP2001460	F-NT2RP200146		
NT2RP2001756	F-NT2RP200175	6 0.17	

NT2RP2002056	F-NT2RP2002056	0. 12
NT2RP2002677	F-NT2RP2002677	0.14
NT2RP2002755	F-NT2RP2002755	0.12
NT2RP2002843	F-NT2RP2002843	0.11
NT2RP2003101	F-NT2RP2003101	0.13
NT2RP2003799	F-NT2RP2003799	0.24
NT2RP2004095	F-NT2RP2004095	0.16
NT2RP2004732	F-NT2RP2004732	0.18
NT2RP2004920	F-NT2RP2004920	0.15
NT2RP2005454	F-NT2RP2005454	0.09
NT2RP2005776	F-NT2RP2005776	0.19
NT2RP2005806	F-NT2RP2005806	0. 27
NT2RP2005882	F-NT2RP2005882	0.11
NT2RP3001282	F-NT2RP3001282	0.39
NT2RP3001723	F-NT2RP3001723	0.22
NT2RP3002099	F-NT2RP3002099	0.20
NT2RP3003155	F-NT2RP3003155	0.29
NT2RP3004028	F-NT2RP3004028	0. 13
0VARC1000008	F-0VARC1000008	0.23
0VARC1000724	F-0VARC1000724	0.27
0VARG1000751	F-0VARC1000751	0. 28
OVARC1001029	F-0VARC1001029	0.25
PLACE1000814	F-PLACE1000814	0. 21
PLACE1003030	F-PLACE1003030	0. 26
PLACE1005549	F-PLACE1005549	0.16
PLACE1007218	F-PLACE1007218	0.30

Table 511

5	clone name	name of the 5'-end	SEQ ID NO	name of the 3'-end	SEQ ID NO of the
		sequence	5' -end	sequence	3' -end
			sequence		sequence
10					
	HEMBA1000497	F-HEMBA10004	97 16111	R-HEMBA100049	7 16165
	HEMBA1001750	F-HEMBA10017	50 16112	R-HEMBA100178	0 16166
	HEMBA1003854	F-HEMBA10038	54 16113	R-HEMBA100385	4 16167
15	HEMBA1004193	F-HEMBA10041	93 16114	R-HEMBA100419	3 16168
	HEMBA1004860	F-HEMBA10048	60 16115	R-HEMBA100486	0 16169
	HEMBA1005572	F-HEMBA10055	72 16116	R-HEMBA100557	72 16170
20	HEMBA1006038	F-HEMBA10060	38 16117	R-HEMBA100603	88 16171
	HEMBA1006092	F-HEMBA10060	92 16118	R-HEMBA100609	2 16172
	HEMBA1006406	F-HEMBA10064	06 16119	R-HEMBA100640	06 16173
	HEMBA1006650	F-HEMBA10066	50 16120	R-HEMBA100665	0 16174
25	HEMBA1006812	F-HEMBA10068	12 16121	R-HEMBA100681	2 16175

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HEMBB1000672 F-HEMBB1000672 16122 R-HEMBB1000672 16176
             HEMBB1001197 F-HEMBB1001197 16123 R-HEMBB1001197 16177
             HEMBB1001871 F-HEMBB1001871 16124 R-HEMBB1001871 16178
             MAMMA1001252 F-MAMMA1001252 16125 R-MAMMA1001252 16179
             MAMMA1002094 F-MAMMA1002094 16126 R-MAMMA1002094 16180
             NT2RM4000634 F-NT2RM4000634 16127 R-NT2RM4000634 16181
             NT2RM4000657 F-NT2RM4000657 16128 R-NT2RM4000657 16182
             NT2RM4000783 F-NT2RM4000783 16129 R-NT2RM4000783 16183
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             NT2RM4000857 F-NT2RM4000857 16130 R-NT2RM4000857 16184
             NT2RM4001178 F-NT2RM4001178 16131 R-NT2RM4001178 16185
             NT2RM4002420 F-NT2RM4002420 16132 R-NT2RM4002420 16186
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             NT2RP2000198 F-NT2RP2000198 16133 R-NT2RP2000198 16187
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             NT2RP2000660 F-NT2RP2000660 16135 R-NT2RP2000660 16189
              NT2RP2001214 F-NT2RP2001214 16136 R-NT2RP2001214 16190
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              NT2RP2001460 F-NT2RP2001460 16137 R-NT2RP2001460 16191
              NT2RP2001756 F-NT2RP2001756 16138 R-NT2RP2001756 16192
              NT2RP2002056 F-NT2RP2002056 16139 R-NT2RP2002056 16193
              NT2RP2002677 F-NT2RP2002677 16140 R-NT2RP2002677 16194
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              NT2RP2002755 F-NT2RP2002755 16141 R-NT2RP2002755 16195
              NT2RP2002843 F-NT2RP2002843 16142 R-NT2RP2002843 16196
              NT2RP2003101 F-NT2RP2003101 16143 R-NT2RP2003101 16197
              NT2RP2003799 F-NT2RP2003799 16144 R-NT2RP2003799 16198
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              NT2RP2004095 F-NT2RP2004095 16145 R-NT2RP2004095 16199
              NT2RP2004732 F-NT2RP2004732 16146 R-NT2RP2004732 16200
              NT2RP2004920 F-NT2RP2004920 16147 R-NT2RP2004920 16201
              NT2RP2005454 F-NT2RP2005454 16148 R-NT2RP2005454 16202
35
              NT2RP2005776 F-NT2RP2005776 16149 R-NT2RP2005776 16203
              NT2RP2005806 F-NT2RP2005806 16150 R-NT2RP2005806 16204
              NT2RP2005882 F-NT2RP2005882 16151 R-NT2RP2005882 16205
              NT2RP3001282 F-NT2RP3001282 16152 R-NT2RP3001282 16206
              NT2RP3001723 F-NT2RP3001723 16153 R-NT2RP3001723 16207
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              NT2RP3002099 F-NT2RP3002099 16154 R-NT2RP3002099 16208
              NT2RP3003155 F-NT2RP3003155 16155 R-NT2RP3003155 16209
              NT2RP3004028 F-NT2RP3004028 16156 R-NT2RP3004028 16210
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              OVARC1000008 F-OVARC1000008 16157 R-OVARC1000008 16211
               OVARC1000724 F-0VARC1000724 16158 R-0VARC1000724 16212
               OVARC1000751 F-0VARC1000751 16159 R-0VARC1000751 16213
              OVARC1001029 F-0VARC1001029 16160 R-0VARC1001029 16214
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              PLACE1000814 F-PLACE1000814 16161 R-PLACE1000814 16215
              PLACE1003030 F-PLACE1003030 16162 R-PLACE1003030 16216
               PLACE1005549 F-PLACE1005549 16163 R-PLACE1005549 16217
               PLACE1007218 F-PLACE1007218 16164 R-PLACE1007218 16218
55
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### **EXAMPLE 17**

Search for a signal sequence, transmembrane region and functional domain in deduced amino acid sequences

- 9 (0552) The deduced amino acid sequences from the full-length nucleotide sequences were examined to predict the presence of a signal sequence in their amino-termin is a well as the presence of a transmembrane region. The amino acid sequences were also searched for a protein functional domain (motif). The examinations for a signal sequence in the amino-terminus, for a transmembrane region and for a functional domain were performed by using PSORT [K. Nakai & M. Kanehisa, Genomics, 14:897-911 (1992)], SOSUI [T. Hirokawa et al., Bioinformatics, 14:378-379 (1998)]
  (Mitsui Knowledge Industry Co., Ltd.) and Pfem (http://www.sanger.ac.uk/Software/Pfaminidex shmin), respectively. When the presence of a signal sequence or a transmembrane region in the amino-terminus was predicted in the amino acid sequence by PSORT or SOSUI, the protein was predicted to be a secretory protein or a membrane protein. When the amino acid sequence matched a functional domain in the Pfam search for a functional domain, the function of the protein is predictable based on the matching data, for example, by referring to the functional categories in PROSITE in (http://www.cpasy.ch/orgi-bin/prosite-list/p). The functional domain search can be performed by using PROSITE in (http://www.cpasy.ch/orgi-bin/prosite-list/p.). The functional domain search can be performed by using PROSITE in 1900 of the protein is provided to the protein in the protein in
  - [0253] Search results obtained by using the respective software programs are indicated below.
    - [0254] Clones whose deduced amino acid sequences were predicted to have signal sequences by PSORT search

```
HEMBA1001052, HEMBA1001407, HEMBA1002486, HEMBA1002661, HEMBA1002818, HEMBA1002876,
HEMBA1003086. HEMBA1003711. HEMBA1004752. HEMBA1005991. HEMBA1006067. HEMBA1006173.
HEMBA1006198, HEMBA1006789, HEMBA1006921, HEMBB1000054, HEMBB1000175, HEMBB1002692,
MAMMA1000798, MAMMA1002427, MAMMA1002881, MAMMA1003035, NT2RM1000035, NT2RM1000742,
NT2RM1000811, NT2RM1000905, NT2RM1001008, NT2RM2000287, NT2RM2000609, NT2RM2001613,
NT2RM4000634, NT2RM4000778, NT2RM4002339, NT2RM4002460, NT2RP1000782, NT2RP1000856,
NT2RP1001247, NT2RP1001546, NT2RP1001569, NT2RP2001597, NT2RP2002537, NT2RP2004142,
NT2RP2005752, NT2RP2005812, NT2RP3001084, NT2RP3001589, NT2RP3002163, NT2RP3002650,
NT2RP3003145, NT2RP3003242, NT2RP3003621, NT2RP3004282, NT2RP3004503, NT2RP4000051,
NT2RP4000151, NT2RP4000243, NT2RP4000259, NT2RP4000323, NT2RP4000417, NT2RP4001064,
NT2RP4001117. NT2RP4001730. NT2RP4001739. NT2RP4002075. NT2RP5003500. OVARC1001154.
PLACE1003031, PLACE1003030, PLACE1003044, PLACE1003369, PLACE1003596, PLACE1004258,
PLACE1005086, PLACE1006239, PLACE1006754, PLACE1006829, PLACE1007954, PLACE1008424,
PLACE1008533. PLACE1008693. PLACE1010622. PLACE1010942. PLACE2000176. PLACE2000341.
PLACE2000379, PLACE2000427, PLACE2000477, PLACE4000431, PLACE4000593, THYRO1000156,
THYRO1001134, THYRO1001287, Y79AA1000258, Y79AA1001874, Y79AA1002399 HEMBB1001871,
HEMBB1001925, MAMMA1000778, MAMMA1000897, MAMMA1001080, NT2RP2004300, NT2RP3002985,
NT2RP3003059, OVARC1000689, OVARC1000890, PLACE1005162, PLACE3000399, PLACE3000455,
PLACE4000247, PLACE4000259, PLACE4000494
```

[0255] Clones whose deduced amino acid sequences were predicted to have transmembrane regions by SOSUI search are as follows:

HEMBA100005, HEMBA1000356, HEMBA1000518, HEMBA1000531, HEMBA1000637, HEMBA1000719, HEMBA1000817, HEMBA1000822, HEMBA1000870, HEMBA1000991, HEMBA1001052, HEMBA1001085, HEMBA1001286. HEMBA1001351. HEMBA1001407. HEMBA1001446. HEMBA1001510. HEMBA1001515. HEMBA1001557, HEMBA1001746, HEMBA1002092, HEMBA1002125, HEMBA1002150, HEMBA1002166, HEMBA1002462. HEMBA1002477. HEMBA1002486. HEMBA1002609. HEMBA1002659. HEMBA1002661. HEMBA1002780, HEMBA1002818, HEMBA1002876, HEMBA1002921, HEMBA1003077, HEMBA1003079, HEMBA1003086, HEMBA1003096, HEMBA1003281, HEMBA1003286, HEMBA1003711, HEMBA1003742, HEMBA1003803, HEMBA1004143, HEMBA1004146, HEMBA1004341, HEMBA1004461, HEMBA1004577, HEMBA1004637, HEMBA1004752, HEMBA1004756, HEMBA1004850, HEMBA1004889, HEMBA1004923, HEMBA1004930, HEMBA1005029, HEMBA1005035, HEMBA1005050, HEMBA1005552, HEMBA1005588, HEMBA1005616, HEMBA1005991, HEMBA1006036, HEMBA1006067, HEMBA1006293, HEMBA1006492, HEMBA1006502, HEMBA1006659, HEMBA1006758, HEMBA1006789, HEMBA1006921, HEMBA1006926, HEMBA1007203, HEMBB1000050, HEMBB1000054, HEMBB1000556, HEMBB1000593, HEMBB1000631, HEMBB1000763. HEMBB1000827. HEMBB1000915. HEMBB1000975. HEMBB1001112. HEMBB1001177. HEMBB1001302. HEMBB1001348. HEMBB1001962. HEMBB1002142. HEMBB1002190. HEMBB1002247. HEMBB1002387, HEMBB1002550, HEMBB1002600, HEMBB1002692, MAMMA1000129, MAMMA1000133, MAMMA1000277, MAMMA1000278, MAMMA1000410, MAMMA1000416, MAMMA1000472, MAMMA1000714, MAMMA1000731, MAMMA1000734, MAMMA1000798, MAMMA1000842, MAMMA1000956, MAMMA1001008,

	MAMMA1001030,	MAMMA1001139,	MAMMA1001154,	MAMMA1001388,	MAMMA1001411,	MAMMA1001487,
	MAMMA1001751,	MAMMA1001771,	MAMMA1002461,	MAMMA1002524,	MAMMA1002598,	MAMMA1002684,
	MAMMA1002769,	MAMMA1002890,	MAMMA1002938,	MAMMA1003146,	NT2RM1000035,	NT2RM1000037,
	NT2RM1000062,	NT2RM1000131,	NT2RM1000257,	NT2RM1000260,	NT2RM1000355,	NT2RM1000648,
5	NT2RM1000742,	NT2RM1000800,	NT2RM1000811,	NT2RM1000857,	NT2RM1000867,	NT2RM1000882,
	NT2RM1001008,	NT2RM1001115,	NT2RM1001139,	NT2RM2000259,	NT2RM2000395,	NT2RM2000402,
	NT2RM2000407,	NT2RM2000422,	NT2RM2000566,	NT2RM2000581,	NT2RM2000609,	NT2RM2001370,
	NT2RM2001393,	NT2RM2001499,	NT2RM2001613,	NT2RM2001648,	NT2RM2001659,	NT2RM2001671,
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	NT2RP2000892,	NT2RP2001081,	NT2RP2001268,	NT2RP2001295,	NT2RP2001366,	NT2RP2001576,
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	PLACE1002213,	PLACE1002395,	PLACE1002500,	PLACE1002714,	PLACE1002722,	PLACE1002794,
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	PLACE1004840,	PLACE1004969,	PLACE1005086,	PLACE1005206,	PLACE1005313,	PLACE1005530,
	PLACE1005595,	PLACE1005623,	PLACE1005763,	PLACE1005884,	PLACE1005934,	PLACE1006225,
	PLACE1006754,	PLACE1006901,	PLACE1006935,	PLACE1006956,	PLACE1007014,	PLACE1007111,

	PLACE1007243,	PLACE1007274,	PLACE1007282,	PLACE1007317,	PLACE1007375,	PLACE1007386,
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	OVARC1001329,	PLACE1001761,	PLACE1002437,	PLACE1004793,	PLACE1005611,	PLACE1005898,
	PLACE1009935,	PLACE1011896,	PLACE2000132,	PLACE2000335,	PLACE3000373,	PLACE3000406,
	PLACE4000250,	PLACE4000487,	PLACE4000494,	THYRO1001320,	THYRO1001537,	THYRO1001828,
	Y79AA1001384					
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[0256] Names of clones whose deduced amino acid sequences were predicted to have functional domains by Pfam search, and names of the matched functional domains are shown below.

When multiple functional domains matched a clone, each domain name was indicated, separated by a double-slash mark. //.

HEMBA1000005//DnaJ, prokaryotic heat shock protein

HEMBA1000020//Tubulin

HEMBA1000129//Helicases conserved C-terminal domain

HEMBA1000156//RNA recognition motif. (aka RRM, RBD, or RNP domain)

HEMBA1000158//Fork head domain, eukaryotic transcription factors //Zinc finger, C2H2 type

35 HEMBA1000303//Src homology domain 3 //Zinc finger, C3HC4 type (RING finger) HEMBA1000411//Ank repeat

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HEMBA1000491//Ras family (contains ATP/GTP binding P-loop)

HEMBA1000531//Heat shock hsp70 proteins

HEMBA1000561//Zinc finger, C2H2 type HEMBA1000608//Src homology domain 3

HEMBA1000919//WD domain, G-beta repeats

HEMBA1001043//Ank repeat

HEMBA1001088//LIM domain containing proteins

HEMBA1001137//Zinc finger, C2H2 type

HEMBA1001174//ADP-ribosylation factors (Arf family) (contains ATP/GTP binding P-loop)

HEMBA1001247//WW/rsp5/WWP domain containing proteins

HEMBA1001286//Sushi domain

HEMBA1001510//Basic region plus leucine zipper transcription factors

HEMBA1001515//Reverse transcriptase (RNA-dependent DNA polymerase)

HEMBA1001661//Cadherin

HEMBA1001723/WD domain. G-beta repeats

HEMBA1001744//Eukaryotic protein kinase domain

HEMBA1001804//Zinc finger, C2H2 type HEMBA1001819//Zinc finger, C2H2 type

55 HEMBA1001847//Zinc finger, C2H2 type

HEMBA1002035//Bromodomain

HEMBA1002102//Ank repeat

HEMBA1002161//Myosin head (motor domain) (contains ATP/GTP binding P-loop)

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HEMBA1002177//GATA family of transcription factors //Zinc finger, C2H2 type HEMBA1002212//Eukaryotic protein kinase domain HEMBA1002215//LIM domain containing proteins HEMBA1002419//RNA recognition motif. (aka RRM, RBD, or RNP domain) HEMBA1002547//Kazal-type serine protease inhibitor domain //Laminin EGF-like (Domains III and V) HEMBA1002768//Src homology domain 3 HEMBA1002810//WW/rsp5/WWP domain containing proteins HEMBA1002818//EGF-like domain HEMBA1002935//Zinc finger, C2H2 type HEMBA1002939//Ank repeat HEMBA1002973//3'5'-cyclic nucleotide phosphodiesterases HEMBA1003077//Fibronectin type III domain HEMBA1003250//Eukaryotic protein kinase domain HEMBA1003257//Zinc finger, C2H2 type HEMBA1003281//IG superfamily HEMBA1003291//Eukarvotic protein kinase domain HEMBA1003433//Forkhead-associated (FHA) domain HEMBA1003545//Homeobox domain //LIM domain containing proteins HEMBA1003591//RNA recognition motif. (aka RRM, RBD, or RNP domain) HEMBA1003684//Zinc finger, C2H2 type HEMBA1003953//Zinc finger, C2H2 type HEMBA1004202//Ras family (contains ATP/GTP binding P-loop) HEMBA1004227//Protein phosphatase 2C HEMBA1004321//Zinc finger, C2H2 type HEMBA1004356//RNA recognition motif. (aka RRM, RBD, or RNP domain) HEMBA1004408//Peptidyl-prolyl cis-trans isomerases HEMBA1004596//RNA recognition motif. (aka RRM, RBD, or RNP domain) HEMBA1004734//Ubiquitin-conjugating enzymes HEMBA1004973//Fibronectin type III domain HEMBA1005009//Actins HEMBA1005101//RNA recognition motif. (aka RRM, RBD, or RNP domain) HEMBA1005581//EGF-like domain //Laminin G domain HEMBA1005732//Polyprenyl synthetases HEMBA1005737//EF hand HEMBA1006248//Zinc finger, C2H2 type HEMBA1006284//Ubiquitin family HEMBA1006293//IG superfamily HEMBA1006344//Band 4.1 family HEMBA1006445//Ras family (contains ATP/GTP binding P-loop) HEMBA1006492//Ank repeat HEMBA1006559//Zinc finger, C3HC4 type (RING finger) HEMBA1006708//WD domain. G-beta repeats HEMBA1006737//Ank repeat HEMBA1006758//Cadherin HEMBA1006941//Thioredoxins HEMBA1007243//Purine/pyrimidine phosphoribosyl transferases HEMBA1007300//3'5'-cyclic nucleotide phosphodiesterases HEMBB1000083/IG superfamily HEMBB1000317//EGF-like domain //Thrombospondin type 1 domain HEMBB1000556//Actinin-type actin-binding domain containing proteins //LIM domain containing proteins HEMBB1000725//Ras family (contains ATP/GTP binding P-loop) HEMBB1000781//Eukarvotic protein kinase domain HEMBB1000915//Thrombospondin type 1 domain HEMBB1000927//EF hand HEMBB1000947//Double-stranded RNA binding motif HEMBB1001112//eubacterial secY protein HEMBB1001175//Ank repeat

HEMBB1001234//WW/rsp5/WWP domain containing proteins

EP 1 074 617 A2 HEMBB1001282//Ank repeat HEMBB1001294//Ras family (contains ATP/GTP binding P-loop) HEMBB1001339//Forkhead-associated (FHA) domain HEMBB1001673//Forkhead-associated (FHA) domain //Zinc finger, C3HC4 type (RING finger) HEMBB1001802//Intermediate filament proteins HEMBB1001839//Zinc finger, C2H2 type HEMBB1002217//Zinc finger, C2H2 type HEMBR1002342//Thioredoxins HEMBB1002600//4 transmembrane segments integral membrane proteins MAMMA1000173//Src homology domain 3 MAMMA1000388//Zinc finger, C2H2 type MAMMA1000402//Reverse transcriptase (RNA-dependent DNA polymerase) MAMMA1000612//WD domain. G-beta repeats MAMMA1000672//Serine carboxypeptidases MAMMA1000731//SNF2 and others N-terminal domain MAMMA1001008//Eukaryotic aspartyl proteases MAMMA1001041//Actinin-type actin-binding domain containing proteins MAMMA1001059//DEAD and DEAH box helicases //Helicases conserved C-terminal domain MAMMA1001105//Zinc finger, C2H2 type MAMMA1001260//Zinc finger, C3HC4 type (RING finger) MAMMA1001576//Tubulin MAMMA1001735//Tubulin MAMMA1001768//ATPases associated with various cellular activities (AAA) MAMMA1001837//Zinc finger, C2H2 type MAMMA1002170//Ribosomal protein S5 MAMMA1002385//RNA recognition motif. (aka RRM, RBD, or RNP domain) MAMMA1002619//Ubiquitin carboxyl-terminal hydrolases family 2 MAMMA1002637//Kinesin light chain repeat MAMMA1002650//Zinc finger, C2H2 type MAMMA1002671//AMP-binding enzymes MAMMA1002869//LIM domain containing proteins MAMMA1002881//SCP-like extracellular Proteins MAMMA1002937//Zinc finger, C2H2 type MAMMA1002938//Multicopper oxidases MAMMA1003011//Core histones H2A, H2B, H3 and H4 MAMMA1003057//WD domain, G-beta repeats MAMMA1003127//Myosin head (motor domain) (contains ATP/GTP binding P-loop) NT2RM1000086//Zinc finger, C3HC4 type (RING finger) NT2RM1000199//CUB domain //Sushi domain NT2RM1000256//Glutamine amidotransferases class-II NT2RM1000499//Ank repeat NT2RM1000555//'Cold-shock' DNA-binding domain containing proteins NT2RM1000666//'Cold-shock' DNA-binding domain containing proteins //Zinc finger, CCHC class NT2RM1000772//WD domain, G-beta repeats NT2RM1000826//'Cold-shock' DNA-binding domain containing proteins NT2RM1000850//Ank repeat //Eukaryotic protein kinase domain NT2RM1000852//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RM1000882//Heme-binding domain in cytochrome b5 and oxidoreductases NT2RM1000885//Zinc finger, C3HC4 type (RING finger) NT2RM1001059//RNA recognition motif. (aka RRM, RBD, or RNP domain)

NT2RM2000092//Ubiquitin carboxyl-terminal hydrolases family 2 NT2RM2000101//Zinc finger, C3HC4 type (RING finger)

NT2RM2000191//3'5'-cyclic nucleotide phosphodiesterases NT2RM2000422//Sodium:neurotransmitter symporter family

specific phospholipase C. Y domain

NT2RM2000490//C2 domain NT2RM2000566//Integrins alpha chain

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NT2RM1001072//C2 domain //Phosphatidylinositol-specific phospholipase C, X domain //Phosphatidylinositol-

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NT2RM4001979//Zinc finger, C2H2 type NT2RM4001987//IG superfamily NT2RM4002013//WD domain. G-beta repeats

NT2RM2000577//tRNA synthetases class I NT2RM2000594//C-5 cytosine-specific DNA methylases NT2RM2000691//Actins NT2RM2000735//Zinc finger, C2H2 type NT2RM2000740//Helicases conserved C-terminal domain NT2RM2000951//FGGY family of carbohydrate kinases NT2RM2001324//LIM domain containing proteins NT2RM2001499//Amino acid permeases NT2RM2001547//DnaJ. prokarvotic heat shock protein //Thioredoxins NT2RM2001613//eubacterial secY protein NT2RM2001670//Zinc finger, C2H2 type NT2RM2001700//Acvl-CoA dehydrogenases NT2RM2001730//Ubiquitin carboxyl-terminal hydrolases family 2 NT2RM2001813//WD domain, G-beta repeats NT2RM2001823//Helicases conserved C-terminal domain //SNF2 and others N-terminal domain NT2RM2001896//Cytochrome C oxidase subunit II NT2RM2001989//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RM2001997//Thioredoxins NT2RM2002088//KH domain family of RNA binding proteins NT2RM2002100//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RM2002109//IG superfamily NT2RM4000046//Zinc finger, C3HC4 type (RING finger) NT2RM4000104//Zinc finger, C2H2 type NT2RM4000167//Kinesin motor domain NT2RM4000191//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RM4000202//Zinc finger, C2H2 type NT2RM4000229//PH (pleckstrin homology) domain NT2RM4000344//ATPases associated with various cellular activities (AAA) NT2RM4000356//Ras family (contains ATP/GTP binding P-loop) NT2RM4000471//Aminotransferases class-V NT2RM4000496//ATPases associated with various cellular activities (AAA) NT2RM4000611//WD domain, G-beta repeats NT2RM4000657//C2 domain //Phosphatidylinositol-specific phospholipase C, Y domain NT2RM4000712//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 NT2RM4000733//Forkhead-associated (FHA) domain NT2RM4000734//Zinc finger, C2H2 type NT2RM4000751//Zinc finger, C2H2 type NT2RM4000795//Carboxylesterases NT2RM4000996//Zinc finger, C2H2 type NT2RM4001054//eubacterial secY protein NT2RM4001140//Homeohox domain NT2RM4001178//DEAD and DEAH box helicases NT2RM4001200//Zinc finger, C2H2 type NT2RM4001313//Phosphatidylinositol 3- and 4-kinases NT2RM4001316//Acvl-CoA dehydrogenases NT2RM4001320//Src homology domain 3 NT2RM4001411//PH (pleckstrin homology) domain //Src homology domain 2 NT2RM4001454//PH (pleckstrin homology) domain NT2RM4001483//Zinc finger, C2H2 type NT2RM4001629//Src homology domain 3 NT2RM4001758//Eukarvotic protein kinase domain NT2RM4001810//Zinc finger, C2H2 type NT2RM4001813//Lectin C-type domain short and long forms NT2RM4001823//Zinc finger, C2H2 type NT2RM4001828//Zinc finger, C2H2 type

EP 1 074 617 A2 NT2RM4002073//AMP-binding enzymes NT2RM4002093//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RM4002145//IG superfamily NT2RM4002287//Fibronectin type III domain NT2RM4002527//WD domain, G-beta repeats NT2RM4002623//tRNA synthetases class II NT2RP1000101//Zinc finger, C2H2 type NT2RP1000202//Ank repeat NT2RP1000272//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RP1000363//PH (pleckstrin homology) domain NT2RP1000376//Ank repeat NT2RP1000470//DEAD and DEAH box helicases NT2RP1000478//Tubulin NT2RP1000522//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 NT2RP1000677//Kazal-type serine protease inhibitor domain NT2RP1000701//WD domain. G-beta repeats NT2RP1000733//Elongation factor Tu family (contains ATP/GTP binding P-loop) NT2RP1000782//4 transmembrane segments integral membrane proteins NT2RP1000833//3'5'-cyclic nucleotide phosphodiesterases NT2RP1000856//4 transmembrane segments integral membrane proteins NT2RP1000947//Ubiquitin-conjugating enzymes NT2RP1000959//60s Acidic ribosomal protein NT2RP1000966//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RP1001033//Tubulin NT2RP1001080//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RP1001177//Core histones H2A H2B H3 and H4 NT2RP1001247//Transforming growth factor beta like domain NT2RP1001294//WD domain. G-beta repeats NT2RP1001302//WD domain, G-beta repeats NT2RP1001313//Heme-binding domain in cytochrome b5 and oxidoreductases NT2RP1001457//WD domain. G-beta repeats NT2RP1001546//4 transmembrane segments integral membrane proteins NT2RP2000008//Zinc finger, C2H2 type NT2RP2000040//C2 domain NT2RP2000045//DnaJ, prokaryotic heat shock protein NT2RP2000054//Zinc finger, C3HC4 type (RING finger) NT2RP2000070//Cadherin NT2RP2000126//Helicases conserved C-terminal domain //SNF2 and others N-terminal domain NT2RP2000153//RNA recognition motif, (aka RRM, RBD, or RNP domain) NT2RP2000224//PH (pleckstrin, homology) domain. NT2RP2000257//Mitochondrial carrier proteins NT2RP2000329//Adenvlate kinases NT2RP2000414//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RP2000448//PH (pleckstrin homology) domain NT2RP2000660//ATPases associated with various cellular activities (AAA) NT2RP2000668//Eukaryotic protein kinase domain NT2RP2000710//tRNA synthetases class II NT2RP2000764//Aminotransferases class-V NT2RP2000842//7 transmembrane receptor (rhodopsin family) NT2RP2000880//Elongation factor Tu family (contains ATP/GTP binding P-loop)

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NT2RP2000931//RNA recognition motif. (aka RRM, RBD, or RNP domain)

NT2RP2000932//Ank repeat NT2RP2001081//C2 domain NT2RP2001174//Zinc finger, C2H2 type NT2RP2001397//Cyclins

NT2RP2001520//Mitochondrial carrier proteins NT2RP2001597//Zinc finger, C3HC4 type (RING finger) NT2RP2001740//Ubiquitin carboxyl-terminal hydrolases family 2

EP 1 074 617 A2 NT2RP2001748//Polyprenyl synthetases NT2RP2001756//Zinc finger, C2H2 type NT2RP2001839//Eukarvotic protein kinase domain NT2RP2001900//Actins NT2RP2001991//Sodium:neurotransmitter symporter family NT2RP2002058//WD domain, G-beta repeats NT2RP2002124//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 NT2RP2002185//Ubiquitin family NT2RP2002208//Zinc finger, C3HC4 type (RING finger) NT2RP2002256//Cytochrome P450 NT2RP2002479//ABC transporters NT2RP2002503//Zinc finger, C2H2 type NT2RP2002520//Ank repeat NT2RP2002591//Zinc finger, C2H2 type NT2RP2002741//Src homology domain 3 NT2RP2002929//WD domain. G-beta repeats NT2RP2002939//Zinc finger, C2H2 type NT2RP2002959//Ubiquitin-conjugating enzymes NT2RP2002980//Ribosomal protein S10 NT2RP2003137//Ubiquitin family NT2RP2003164//Eukarvotic protein kinase domain NT2RP2003228//MCM2/3/5 family NT2RP2003243//Fibronectin type III domain NT2RP2003272//Ubiquitin family NT2RP2003307//Kinesin light chain repeat NT2RP2003401//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases, family 2 NT2RP2003433//eubacterial secY protein NT2RP2003480//Zinc finger, C2H2 type NT2RP2003713//Ubiquitin carboxyl-terminal hydrolases family 2 NT2RP2003737//Ubiquitin-conjugating enzymes NT2RP2003777//Zinc finger, C3HC4 type (RING finger) NT2RP2003840//Ubiquitin-conjugating enzymes NT2RP2003857//Ank repeat NT2RP2003981//Zinc finger, C3HC4 type (RING finger) NT2RP2004170//WD domain, G-beta repeats NT2RP2004187//Zinc finger, C2H2 type NT2RP2004232//Phorbol esters / diacylglycerol binding domain //PH (pleckstrin homology) domain //Eukaryotic protein kinase domain NT2RP2004389//Ribosomal protein S9 NT2RP2004538//PH (pleckstrin homology) domain NT2RP2004568//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RP2004710//WW/rsp5/WWP domain containing proteins NT2RP2004768//Eukarvotic protein kinase domain NT2RP2004933//Eukaryotic protein kinase domain NT2RP2004961//Zinc finger, C2H2 type NT2RP2005003//Zinc finger, C3HC4 type (RING finger) NT2RP2005012//DnaJ, prokaryotic heat shock protein NT2RP2005126//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RP2005139//Ank repeat NT2RP2005140//PH (pleckstrin homology) domain NT2RP2005239//Aminotransferases class-V

NT2RP2005344//E1-E2 ATPases NT2RP2005525//Forkhead-associated (FHA) domain

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NT2RP2005465//Mitochondrial carrier proteins

NT2RP2005288//Regulator of chromosome condensation (RCC1) NT2RP2005293//PH (pleckstrin homology) domain NT2RP2005325//Homeobox domain //LIM domain containing proteins

NT2RP2005531//Band 4.1 family

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NT2RP2005557//Bacterial mutT protein NT2RP2005654//DnaJ, prokaryotic heat shock protein NT2RP2005701//Zinc finger, C3HC4 type (RING finger) NT2RP2005722//Zinc finger, C2H2 type NT2RP2005752//TNFR/NGFR cysteine-rich region NT2RP2005763//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RP2005767//HMG (high mobility group) box NT2RP2006312//HMG (high mobility group) box NT2RP2006464//HMG (high mobility group) box NT2RP2006571//Cytochrome P450 NT2RP3000050//Zinc finger, C2H2 type NT2RP3000068//PH (pleckstrin homology) domain NT2RP3000085//Biotin-requiring enzymes //Carbamoyl-phosphate synthase (CPSase) NT2RP3000299//Src homology domain 3 NT2RP3000359//Adenylate kinases NT2RP3000366//Ras family (contains ATP/GTP binding P-loop) NT2RP3000403//WW/rsp5/WWP domain containing proteins NT2RP3000487//WW/rsp5/WWP domain containing proteins NT2RP3000512//Homeobox domain NT2RP3000527//Zinc finger, C2H2 type NT2RP3000531//IG superfamily NT2RP3000590//Zinc finger, C3HC4 type (RING finger) NT2RP3000603//Helix-loop-helix DNA-binding domain NT2RP3000605//Zinc finger, C2H2 type NT2RP3000632//Zinc finger, C2H2 type NT2RP3000742//Phosphatidylinositol-specific phospholipase C, X domain //Phosphatidylinositol-specific phospholipase C. Y domain NT2RP3000759//ADP-ribosylation factors (Arf family) (contains ATP/GTP binding P-loop) NT2RP3000825//EGF-like domain NT2RP3000869//ATPases associated with various cellular activities (AAA) NT2RP3000994//Double-stranded RNA binding motif NT2RP3001057//Zinc finger, C2H2 type NT2RP3001084//PH (pleckstrin homology) domain NT2RP3001120//Zinc finger, C2H2 type NT2RP3001140//Thrombospondin type 1 domain NT2RP3001150//Forkhead-associated (FHA) domain NT2RP3001155//HMG (high mobility group) box NT2RP3001214//Zinc finger, C2H2 type NT2RP3001268//Zinc finger, C2H2 type NT2RP3001338//Zinc finger, C2H2 type NT2RP3001355//Mitochondrial carrier proteins NT2RP3001398//Zinc finger, C2H2 type NT2RP3001426//DnaJ, prokaryotic heat shock protein NT2RP3001453//ABC transporters NT2RP3001457//PH (pleckstrin homology) domain NT2RP3001472//HMG (high mobility group) box NT2RP3001495//Alcohol/other dehydrogenases, short chain type //WW/rsp5/WWP domain containing proteins NT2RP3001497//Zinc finger, C3HC4 type (RING finger) NT2RP3001724//Helicases conserved C-terminal domain NT2RP3001792//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RP3001943//Zinc finger, C3HC4 type (RING finger) NT2RP3001944//Zinc finger, C3HC4 type (RING finger) NT2RP3002007//ATPases associated with various cellular activities (AAA) NT2RP3002054//Low-density lipoprotein receptor domain class A NT2RP3002151//Elongation factor Tu family (contains ATP/GTP binding P-loop) NT2RP3002399//MCM2/3/5 family NT2RP3002501//Serine/threonine dehydratases NT2RP3002602//Thioredoxins

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NT2RP4001498//Ank repeat NT2RP4001568//Ank repeat

NT2RP4001644//Eukaryotic protein kinase domain NT2RP4001725//WD domain, G-beta repeats NT2RP4001753//Zinc finger, C2H2 type NT2RP4001790//Zinc finger, C2H2 type

NT2RP3002628//DnaJ. prokaryotic heat shock protein //Thioredoxins NT2RP3002663//PH (pleckstrin homology) domain NT2RP3002909//Ank repeat NT2RP3002953//Cadherin NT2RP3002969//AMP-binding enzymes NT2RP3003061//Ank repeat NT2RP3003145//Zinc carboxypeptidases NT2RP3003230//WD domain, G-beta repeats NT2RP3003251//Zinc finger, C3HC4 type (RING finger) NT2RP3003278//Ank repeat //Zinc finger, C2H2 type NT2RP3003282//PH (pleckstrin homology) domain NT2RP3003311//PH (pleckstrin homology) domain NT2RP3003385//Ank repeat //Chaperonins clpA/B NT2RP3003589//Ras family (contains ATP/GTP binding P-loop) NT2RP3003621//CUB domain //Kringle domain NT2RP3003701//Thrombospondin type 1 domain NT2RP3003716//Fibronectin type III domain NT2RP3003809//ATPases associated with various cellular activities (AAA) NT2RP3004016//Zinc finger, C3HC4 type (RING finger) NT2RP3004207//CUB domain //Sushi domain NT2RP3004209//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 NT2RP3004242//PH (pleckstrin homology) domain NT2RP3004262//DnaJ. prokaryotic heat shock protein NT2RP3004566//Zinc finger, C2H2 type NT2RP3004569//Ank repeat NT2RP3004594//HMG (high mobility group) box NT2RP3004617//Zinc finger, C3HC4 type (RING finger) NT2RP4000259//Glutathione peroxidases NT2RP4000370//Prokaryotic-type class I peptide chain release factors NT2RP4000376//WD domain. G-beta repeats NT2RP4000398//Zinc finger, C2H2 type NT2RP4000455//Forkhead-associated (FHA) domain //Zinc finger, C3HC4 type (RING finger) NT2RP4000457//Ubiquitin carboxyl-terminal hydrolases family 2 NT2RP4000518//DEAD and DEAH box helicases //Helicases conserved C-terminal domain NT2RP4000588//Actinin-type actin-binding domain containing proteins NT2RP4000614//RNA recognition motif, (aka RRM, RBD, or RNP domain) NT2RP4000648//Forkhead-associated (FHA) domain //Zinc finger, C3HC4 type (RING finger) NT2RP4000837//Zinc finger, C2H2 type NT2RP4000839//WD domain, G-beta repeats NT2RP4000865//Zinc finger, C2H2 type NT2RP4000907//Fibronectin type III domain //IG superfamily NT2RP4000925//Fibronectin type III domain NT2RP4000927//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 NT2RP4000973//DnaJ, prokaryotic heat shock protein //Thioredoxins NT2RP4001079//E1-E2 ATPases NT2RP4001080//RNA recognition motif. (aka RRM, RBD, or RNP domain) NT2RP4001117//eubacterial secY protein NT2RP4001150//Fibronectin type III domain NT2RP4001213//Zinc finger, C2H2 type NT2RP4001219//Thioredoxins NT2RP4001235//Zinc finger, CCHC class NT2RP4001433//Zinc finger, C2H2 type

NT2RP4001822//4 transmembrane segments integral membrane proteins NT2RP4001823//Fibrinogen beta and gamma chains, C-terminal globular domain NT2RP4001893//Ank repeat NT2RP4001896//WD domain. G-beta repeats NT2RP4001927//WD domain, G-beta repeats NT2RP4001938//Zinc finger, C2H2 type NT2RP4002047//Elongation factor Tu family (contains ATP/GTP binding P-loop) NT2RP4002078//Zinc finger, C2H2 type NT2RP4002408//Eukarvotic protein kinase domain NT2RP4002905//Cyclins NT2RP5003477//WD domain, G-beta repeats OVARC1000006//Core histones H2A, H2B, H3 and H4 OVARC1000085//Proteasome A-type and B-type OVARC1000148//RNA recognition motif. (aka RRM, RBD, or RNP domain). OVARC1000556//Eukarvotic protein kinase domain OVARC1000649//PH (pleckstrin homology) domain //Src homology domain 2 OVARC1000746//Double-stranded RNA binding motif OVARC1000885//Alcohol/other dehydrogenases, short chain type OVARC1000937//Cyclins OVARC1000999//Ank reneat OVARC1001154//Granulins OVARC1001180//Ubiquitin family OVARC1001306//Helix-loop-helix DNA-binding domain OVARC1001577//RNA recognition motif. (aka RRM, RBD, or RNP domain). OVARC1001731//Tropomyosins OVARC1001943//Zinc finger, C2H2 type OVARC1002050//Spectrin alpha chain, repeated domain OVARC1002112//Core histones H2A, H2B, H3 and H4 OVARC1002138//ATPases associated with various cellular activities (AAA) OVARC1002182//WD domain. G-beta repeats PLACE1000014//Zinc finger, C3HC4 type (RING finger) PLACE1000040//Ras family (contains ATP/GTP binding P-loop) PLACE 1000050//Zinc finger, C2H2 type PLACE1000081//PH (pleckstrin homology) domain PLACE1000142//Enovi-CoA hydratase/isomerase PLACE1000401//IG superfamily PLACE1000406//RNA recognition motif. (aka RRM, RBD, or RNP domain) PLACE1000420//Bacterial mutT protein PLACE1000706//Bromodomain PLACE1000769//KH domain family of RNA binding proteins PLACE1000786//PH (pleckstrin homology) domain PLACE1000863//Ribosomal protein S4 PLACE1000909//Ank repeat PLACE1000972//Src homology domain 3 PLACE1000979//Zinc finger, C2H2 type PLACE1001304//Zinc finger, C2H2 type PLACE1001387//Src homology domain 3 PLACE 1001632//Zinc finger, C2H2 type PLACE1001672//Aminotransferases class-III pyridoxal-phosphate PLACE1001716//Zinc finger, CCHC class PLACE1001739//DEAD and DEAH box helicases //Helicases conserved C-terminal domain PLACE1001781//Phosphoglucomutase and phosphomannomutase phosphoserine PLACE1001869//FGGY family of carbohydrate kinases PLACE1002438//Zinc finger, C2H2 type PLACE 1002450//Zinc finger, C2H2 type PLACE1002474//EGF-like domain //von Willebrand factor type A domain

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PLACE1002499//Zinc finger, C3HC4 type (RING finger) PLACE1002532//Homeobox domain

EP 1 074 617 A2 PLACE1002571//Actins PLACE1002685//Src homology domain 2 PLACE1002722//7 transmembrane receptor (rhodopsin family) PLACE1002775//Bromodomain PLACE1002834//Zinc finger, C2H2 type PLACE1003100//Alcohol/other dehydrogenases, short chain type PLACE1003174//Ubiquitin-conjugating enzymes PLACE1003238//7 transmembrane receptor (rhodopsin family) PLACE1003302//Zinc finger, C2H2 type PLACE1003334//RNA recognition motif. (aka RRM, RBD, or RNP domain) PLACE1003366//C2 domain PLACE1003394//Ras family (contains ATP/GTP binding P-loop) PLACE1003420//Mitochondrial carrier proteins PLACE1003493//C1g domain PLACE1003519//KH domain family of RNA binding-proteins PLACE1003723//Src homology domain 2 PLACE1003738//Zinc finger, C2H2 type PLACE1003888//C2 domain //Phosphatidylinositol-specific phospholipase C, X domain //Phosphatidylinositolspecific phospholipase C, Y domain PLACE1004128//WD domain, G-beta repeats PLACE1004358//PH (pleckstrin homology) domain PLACE1004428//Acyl-CoA dehydrogenases PLACE1004437//isocitrate and isopropylmalate dehydrogenases PLACE1004506//LIM domain containing proteins PLACE1004674//EF hand PLACE1004918//L-lactate dehydrogenases PLACE1005243//Eukarvotic protein kinase domain PLACE1005305//Adenylate kinases PLACE1005327//Src homology domain 3 PLACE1005530//Zinc finger, C3HC4 type (RING finger) PLACE1005646//Helicases conserved C-terminal domain PLACE1005656//Ribonucleotide reductases PLACE1005966//WD domain, G-beta repeats PLACE1006157//Sushi domain PLACE1006196//DEAH and DEAR box helicases //Helicases conserved C-terminal domain PLACE1006438//Zinc finger, C2H2 type PLACE1006626//Double-stranded RNA binding motif PLACE1006754//IG superfamily PLACE1006829//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 PLACE1006917//RNA recognition motif. (aka RRM, RBD, or RNP domain) PLACE1006956//ABC transporters PLACE1006958//Heat shock hsp70 proteins PLACE1007375//C2 domain PLACE1007488//PH (pleckstrin homology) domain PLACE1007511//Intermediate filament proteins PLACE1007537//Ank repeat PLACE1007544//Zinc finger, C2H2 type PLACE1007547//Zinc finger, C3HC4 type (RING finger) PLACE1007598//Zinc finger, C2H2 type PLACE1007697//ABC transporters PLACE1007958//3'5'-cyclic nucleotide phosphodiesterases

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PLACE1007969//RNA recognition motif. (aka RRM, RBD, or RNP domain)

PLACE1008201//Zinc finger, C2H2 type PLACE1008429//Ank repeat PLACE1008465//Zinc finger, C2H2 type PLACE1008650//WD domain. G-beta repeats PLACE1009020//Aminotransferases class-V PLACE1009094//von Willebrand factor type C domain

EP 1 074 617 A2 PLACE1009099//Zinc finger, C2H2 type PLACE1009246//LIM domain containing proteins PLACE1009468//WD domain, G-beta repeats PLACE1009476//DEAD and DEAH box helicases //Helicases conserved C-terminal domain PLACE1009524//PH (pleckstrin homology) domain PLACE1009596//WD domain, G-beta repeats PLACE1009622//Double-stranded RNA binding motif PLACE1009861//Cysteine proteases PLACE1009925//Helicases conserved C-terminal domain PLACE1009992//CUB domain //EGF-like domain //Sushi domain //Trypsin PLACE1010053//Double-stranded RNA binding motif PLACE1010089//Ubiquitin carboxyl-terminal hydrolases family 2 PLACE1010702//Zinc finger, C2H2 type PLACE1010833//EF hand PLACE1010926//Src homology domain 3 PLACE1010960//Actins PLACE1011041//Src homology domain 3 PLACE1011046//C2 domain //Phosphatidylinositol-specific phospholipase C, X domain //Phosphatidylinositol-specific phospholipase C. Y domain PLACE1011114//Helicases conserved C-terminal domain. PLACE1011160//Zinc finger, C3HC4 type (RING finger) PLACE1011263//Ank repeat PLACE1011433//Zinc finger, C2H2 type PLACE1011576//Zinc finger, C2H2 type PLACE1011923//Eukarvotic protein kinase domain PLACE2000034//Fibronectin type III domain //IG superfamily PLACE2000072//Zinc finger, C2H2 type PLACE2000111//IG superfamily PLACE2000164//WD domain, G-beta repeats PLACE2000216//PH (pleckstrin homology) domain PLACE2000341//Sodium:solute symporter family PLACE2000371//Src homology domain 2 PLACE2000373//Thrombospondin type 1 domain PLACE2000398//IG superfamily PLACE2000427//Helicases conserved C-terminal domain PLACE2000458//Cadherin PLACE3000020//Guanviate cyclases PLACE3000169//Zinc finger, C2H2 type PLACE4000014//Helicases conserved C-terminal domain PLACE4000052//ABC transporters PLACE4000192//Zinc finger, C2H2 type PLACE4000211//Bromodomain PLACE4000431//Helicases conserved C-terminal domain PLACE4000522//Ank repeat PLACE4000581//EGF-like domain //Sushi domain PLACE4000654//Ubiquitin-conjugating enzymes THYRO1000072//IG superfamily THYRO1000242//Zinc finger, C2H2 type THYRO1000288//Zinc-binding metalloprotease domain THYRO1000488//Zinc finger, C3HC4 type (RING finger) THYRO1000501//Zinc finger, C3HC4 type (RING finger) THYRO1000666//Kinesin motor domain THYRO1000748//Src homology domain 3 THYRO1000926//3' 5'-cyclic nucleotide phosphodiesterases

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THYRO1001661//RNA recognition motif. (aka RRM, RBD, or RNP domain)

Y79AA1000037//Zinc finger, C3HC4 type (RING finger) Y79AA1000214//Core histones H2A, H2B, H3 and H4

THYRO1001671//Ubiquitin family

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Y79AA1000342//Zinc finger, C2H2 type
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Y79AA1000349//Double-stranded RNA binding motif

Y79AA1000627//Zinc finger, C2H2 type

Y79AA1000705//Helicases conserved C-terminal domain

Y79AA1000752//KH domain family of RNA binding proteins

Y79AA1000833//Tubulin

Y79AA1001048//Acyl-CoA dehydrogenases

Y79AA1001391//Homeobox domain

Y79AA1001394//ATPases associated with various cellular activities (AAA)

Y79AA1001493//Ubiquitin-conjugating enzymes

Y79AA1001613//Zinc finger, C2H2 type

Y79AA1001874//TNFR/NGFR cysteine-rich region

Y79AA1002027//Ubiquitin-conjugating enzymes

Y79AA1002139//DnaJ, prokaryotic heat shock protein Y79AA1002208//Ank repeat

Y79AA1002246//C2 domain

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Y79AA1002307//Fibronectin type III domain

Y79AA1002472//Zinc finger, C2H2 type

HEMBA1003538//CUB domain HEMBA1003645//WD domain, G-beta repeats //Src homology domain 3 HEMBA1005206//Glutathione S-transferases

HEMBA1006521//Alcohol/other dehydrogenases, short chain type

HEMBB1001482//Zinc finger, C2H2 type HEMBB1001915//Ubiquitin carboxyl-terminal hydrolases family 2 //Ubiquitin carboxyl-terminal hydrolases family 2 HEMBB1002044//Cadherin MAMMA1000183//Zinc finger, C2H2 type MAMMA100089//yon Willebrand factor type 4 domain MAMMA100180//G superfamily MAMMA100280//

25 superfamily MAMMA1002573//KH domain family of RNA binding proteins MAMMA1002617//Zincfinger, C2H2 type NT2RM1000833//eubacterial secY protein NT2RM2001797//Zinc finger, C2H2 type

NT2RP1001013//Zinc finger, C2H2 type NT2RP2001233//Zinc finger, C2H2 type

NT2RP2001440//14-3-3 proteins NT2RP2002105//7 transmembrane receptor (rhodopsin family) NT2RP3001723//Laminin G domain NT2RP3001938//Eukaryotic protein kinase domain NT2RP3002330//Elonga-

30 tion factor Tu family (contains ATP/GTP binding P-loop) NT2RP3003133//Zinc finger, C2H2 type NT2RP3003500//Eukaryotic protein kinase domain NT2RP3003799//C2 domain

NT2RP3003800//Eukaryotic protein kinase domain NT2RP3004013//Double-stranded RNA binding motif NT2RP3004125//Zinc finger. C2H2 type

OVARC1001244//Bromodomain OVARC1001496//D-isomer specific 2-hydroxyacid dehydrogenases PLACE1000007/I/bliquitin carboxyl-terminal hydrolases family 2 //l/bliquitin carboxyl-terminal hydrolases family 2 PLACE1001118//Zinc finger, C2H2 type PLACE1010310//Zinc finger, C2H2 type PLACE1011896//wnt family of developmental signaling proteins PLACE3000124//Src homology domain 2

PLACE4000100//D-isomer specific 2-hydroxyacid dehydrogenases

PLACE4000259/I/Helicases conserved C-terminal domain PLACE4000261/I/Bromodomain SKNMC1000013/I/BCD transporters SKNMC10000091/I/Bast region plus leutione zipper transportipion factors THYRO1000343/I/Src homology domain 3 THYRO1000369/I/Zinc finger, C2H2 kpp THYRO1001189/I/Zinc finger, C2H2 kpp Y39AA1002103/I/Zinc finger, C2H2 to PLACE3000350/I/Clack Trottor protain kinase domain

PLACE4000156//Zinc finger, C2H2 type

### 45 EXAMPLE 18

Classification of cDNA clones into functional categories based on the full-length nucleotide sequences

[0257] Prediction of functions of proteins encoded by the clones and the categorization thereof were performed based on the results of homology search (see Homology search results 6, 12, 13 and 14) of the databases, GenBank, Swiss-Prot and UniGene, for the full-length nucleotide sequences of 4997 clones and based on the results of domain search (see Example 17) of the deduced amino acid sequences encoded by the full-length nucleotide sequences. The target 4997 clones are listed below.

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	THYRO1000843,	THYRO1000852,	THYRO1000855,	THYRO1000865,	THYRO1000895,	THYRO1000916,
	THYRO1000926,	THYRO1000934,	THYRO1000951,	THYRO1000952,	THYRO1000983,	THYRO1000988,
	THYRO1001003,	THYRO1001031,	THYRO1001033,	THYRO1001062,	THYRO1001100,	THYRO1001120,
45	THYRO1001133,	THYRO1001134,	THYRO1001142,	THYRO1001173,	THYRO1001189,	THYRO1001204,
45	THYRO1001213,	THYRO1001262,	THYRO1001271,	THYRO1001287,	THYRO1001313,	THYRO1001320,
	THYRO1001321,	THYRO1001322,	THYRO1001347,	THYRO1001363,	THYRO1001365,	THYRO1001374,
	THYRO1001401,	THYRO1001403,	THYRO1001405,	THYRO1001406,	THYRO1001411,	THYRO1001426,
	THYRO1001434,	THYRO1001458,	THYRO1001480,	THYRO1001487,	THYRO1001534,	THYRO1001537,
	THYRO1001541,	THYRO1001559,	THYRO1001570,	THYRO1001584,	THYRO1001595,	THYRO1001602,
50	THYRO1001605,	THYRO1001617,	THYRO1001637,	THYRO1001656,	THYRO1001661,	THYRO1001671,
	THYRO1001673,	THYRO1001703,	THYRO1001706,	THYRO1001721,	THYRO1001738,	THYRO1001745,
	THYRO1001746,	THYRO1001772,	THYRO1001793,	THYRO1001809,	THYRO1001828,	THYRO1001854,
	THYRO1001895,	THYRO1001907,	VESEN1000122,		Y79AA1000033,	Y79AA1000037,
	Y79AA1000059,	Y79AA1000065,	Y79AA1000131,	Y79AA1000181,	Y79AA1000202,	Y79AA1000214,
55	Y79AA1000230,	Y79AA1000231,	Y79AA1000258,	Y79AA1000268,	Y79AA1000313,	Y79AA1000328,
	Y79AA1000342,	Y79AA1000346,	Y79AA1000349,	Y79AA1000355,	Y79AA1000368,	Y79AA1000410,
	Y79AA1000420,	Y79AA1000469,	Y79AA1000480,	Y79AA1000539,	Y79AA1000540,	Y79AA1000560,
	Y79AA1000574,	Y79AA1000589,	Y79AA1000627,	Y79AA1000705,	Y79AA1000734,	Y79AA1000748,

	Y79AA1000752,	Y79AA1000774,	Y79AA1000782,	Y79AA1000784,	Y79AA1000794,	Y79AA1000800,
	Y79AA1000802,	Y79AA1000805,	Y79AA1000824,	Y79AA1000827,	Y79AA1000833,	Y79AA1000850,
	Y79AA1000962,	Y79AA1000966,	Y79AA1000968,	Y79AA1000969,	Y79AA1000976,	Y79AA1000985,
	Y79AA1001023,	Y79AA1001041,	Y79AA1001048,	Y79AA1001061,	Y79AA1001068,	Y79AA1001077,
5	Y79AA1001078,	Y79AA1001145,	Y79AA1001167,	Y79AA1001177,	Y79AA1001185,	Y79AA1001211,
	Y79AA1001216,	Y79AA1001228,	Y79AA1001233,	Y79AA1001236,	Y79AA1001281,	Y79AA1001299,
	Y79AA1001312,	Y79AA1001323,	Y79AA1001384,	Y79AA1001391,	Y79AA1001394,	Y79AA1001402,
	Y79AA1001493,	Y79AA1001511,	Y79AA1001533,	Y79AA1001541,	Y79AA1001548,	Y79AA1001555,
	Y79AA1001581,	Y79AA1001585,	Y79AA1001594,	Y79AA1001603,	Y79AA1001613,	Y79AA1001647,
10	Y79AA1001665,	Y79AA1001679,	Y79AA1001692,	Y79AA1001696,	Y79AA1001705,	Y79AA1001711,
	Y79AA1001781,	Y79AA1001805,	Y79AA1001827,	Y79AA1001846,	Y79AA1001866,	Y79AA1001874,
	Y79AA1001875,	Y79AA1001923,	Y79AA1001963,	Y79AA1002027,	Y79AA1002083,	Y79AA1002089,
	Y79AA1002103,	Y79AA1002115,	Y79AA1002125,	Y79AA1002139,	Y79AA1002204,	Y79AA1002208,
	Y79AA1002209,	Y79AA1002210,	Y79AA1002211,	Y79AA1002220,	Y79AA1002229,	Y79AA1002234,
15	Y79AA1002246,	Y79AA1002258,	Y79AA1002298,	Y79AA1002307,	Y79AA1002311,	Y79AA1002351,
	Y79AA1002361,	Y79AA1002399,	Y79AA1002407,	Y79AA1002416,	Y79AA1002431,	Y79AA1002433,
	Y79AA1002472, Y	779AA1002482, Y79	AA1002487.			

[IQ258] Among the 4997 clones, there are 2189 clones that presumably encode proteins belonging to any of the categories of secretory or membrane proteins, glycoprotein-associated proteins, signal transuclicin-associated proteins, and the categories of secretory or membrane proteins, glycoproteins, enzymes and/or metabolism-associated proteins, ATP- and/or CTP-binding proteins, nuclear proteins, DNA- and/or RNA-binding proteins, RNA synthesis-associated proteins, protein synthesis-and/or protein transport-associated proteins, cytoskeleton-associated proteins, and with the control of th

[0259] The clones that presumably encode proteins belonging to the category of secretory or membrane proteins are those which matched the full-length sequences of Swiss-Prot database with the keywords "growth factor", "cytokine", "hormone", "signal", "transmembrane", "membrane", "cateclalum matrit," receptor, "G-protein coupled enceptor, "lonic channel", "voltage-gated channel", "calcium channel", "cell adhesion", "collagen", or "connective tissue"; those which matched the data, suggesting that the proteins are secretory or membrane proteins, or those which matched the full-length sequences of GenBank or UniGene database with similar description, and, further, those predicted to have an N-terminal signal sequence or a transmembrane region as a result of domain search for the amino acid sequences deduced from the full-length succession.

[0260] The clones that presumably encode proteins belonging to the category of glycoprotein-associated proteins are those which matched the full-length sequences of Swiss-Prot database with the keywords "glycoprotein"; those which matched the data, suggesting that the proteins are glycoprotein; or those which matched the full-length sequences of GenBank or JuriGene database with similar described.

[0261] The clones that presumably encode proteins belonging to the category of signal transduction-associated proteins are those which matched the full-length sequences of Swiss-Prot database with the keywords "serine/threonineprotein kinase"; "tyrosine-protein kinase", or "SH3 domain"; those which matched the data, suggesting that the proteins are signal transduction-associated proteins (for example, "ADP-ribosylation factor"); or those which matched the fulllength sequences of GenBank or UniGene database with similar descriptions.

[0262] The clones that presumably encode proteins belonging to the category of transcription-associated proteins are those which matched the full-length sequences of Swiss-Prot database with the keywords "transcription regulation", "zinc finger", or "homeobox"; those which matched the data, suggesting that the proteins are transcription-associated proteins; or those which matched the full-length sequences of GenBank or UniGene database with similar description. [0263] The clones that presumably encode proteins belonging to the category of disease-associated proteins are those which matched the full-length sequences of Swiss-Prot database with the keywords "disease mutation" or "syn-drome"; those which matched the data, suggesting that the proteins are disease-associated proteins; or those which matched the full-length sequences of Swiss-Prot database and GenBank or UniGene database where the matched sequences of genes or proteins which had been registered in the database of Online Mendellan Inheritance in Man (OMM) (httb://www.ncbi.im.ni.nov/Omirn/), which is a database of human genes and diseasens and series and sequences and severe the matched sequences and sequences are sequences and sequences and sequences and sequences and sequences are sequences and sequences and sequences and sequences are sequences and sequences and sequences and sequences are sequences and sequences are sequences and sequences and sequences are sequences and sequences are sequences and sequences are

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[0264] The clones that presumably encode proteins belonging to the category of enzymes and/or metabolism-associated proteins are those which showed the terms "metabolism", "oxidoreductase", or "E.C. No. (Enzyme commission number)" in the matching data.

[0265] The clones that presumably encode proteins belonging to the category of ATP- and/or GTP-binding proteins are those which matched the data with the terms "ATP-binding" or "GTP-binding".

[0266] The clones that presumably encode proteins belonging to the category of nuclear proteins are those which matched the data with the terms "nuclear protein".

- [0267] The clones that presumably encode proteins belonging to the category of DNA- and/or RNA-binding proteins are those which matched the data with the terms "DNA-binding" or "RNA-binding".
- [0268] The clones that presumably encode proteins belonging to the category of RNA synthesis-associated proteins are those which matched the data with the terms "RNA splicing", "RNA processing", "RNA helicase", or "polyadenylation".
- [0269] The clones that presumably encode proteins belonging to the category of protein synthesis- and/or protein transport-associated proteins are those which matched the data with the terms "translation regulation", "protein biosynthesis," intoxomal protein", "protein bransport", or "sunal recognition particles."
- [0270] The clones that presumably encode proteins belonging to the category of cytoskeleton-associated proteins are those which matched the data with the terms "structural protein", "cytoskeleton", "actin-binding", or "microtubules". [0271] The clones that presumably encode proteins belonging to the category of cell division- and/or cell profileration-associated proteins are those which matched the data with the terms "cell division", "cell cycle", "mitosis", "chromosomal protein", "cell growth", or "apoptosis".
- [0272] The clones that presumably encode proteins belonging to the category of embryogenesis- and/or development-associated proteins are those which matched the data with the terms "developmental protein".
- [0273] The clones that presumably encode proteins belonging to the category of cellular defense-associated proteins are those which matched the data with the terms "heat shock", "DNA repair", or "DNA damage".
- [0274] When a clone belonged to the above-mentioned multiple functional categories, the clone was classified into the multiple categories. However, the functions of the protein encoded by the done are not limited to the functions of the categories into which the clone was classified, and therefore, additional functions can be found for the protein by further analyses.
  - [0275] The following 796 clones are categorized into secretory or membrane proteins.
- HEMBA1000356, HEMBA1000518, HEMBA1000531, HEMBA1000637, HEMBA1000719, HEMBA1000817, HEMBA1000822, HEMBA1000852, HEMBA1000870, HEMBA1000991, HEMBA1001052, HEMBA1001071, HEMBA1001085, HEMBA1001286, HEMBA1001351, HEMBA1001407, HEMBA1001446, HEMBA1001515, HEMBA1001557, HEMBA1001569, HEMBA1001661, HEMBA1001734, HEMBA1001746, HEMBA1001866, HEMBA1002125, HEMBA1002150, HEMBA1002166, HEMBA1002417, HEMBA1002462, HEMBA1002475, HEMBA1002477, HEMBA1002486, HEMBA1002609, HEMBA1002659, HEMBA1002661, HEMBA1002780, HEMBA1002818, HEMBA1002876, HEMBA1002921, HEMBA1003071, HEMBA1003077, HEMBA1003079, HEMBA1003086. HEMBA1003096. HEMBA1003281. HEMBA1003286. HEMBA1003538. HEMBA1003711. HEMBA1003742, HEMBA1003803, HEMBA1004055, HEMBA1004143, HEMBA1004146, HEMBA1004207, HEMBA1004341, HEMBA1004461, HEMBA1004577, HEMBA1004637, HEMBA1004752, HEMBA1004756, HEMBA1004850. HEMBA1004889. HEMBA1004923. HEMBA1004930. HEMBA1005029. HEMBA1005035. HEMBA1005050, HEMBA1005552, HEMBA1005576, HEMBA1005581, HEMBA1005588, HEMBA1005616, HEMBA1005699, HEMBA1005991, HEMBA1006036, HEMBA1006038, HEMBA1006067, HEMBA1006173, HEMBA1006198. HEMBA1006293. HEMBA1006310. HEMBA1006492. HEMBA1006502. HEMBA1006583. HEMBA1006659, HEMBA1006758, HEMBA1006789, HEMBA1006921, HEMBA1006926, HEMBA1006976, HEMBA1007203, HEMBA1007301, HEMBB1000037, HEMBB1000050, HEMBB1000054, HEMBB1000175, HEMBB1000317. HEMBB1000556, HEMBB1000593, HEMBB1000631, HEMBB1000763, HEMBB1000827, HEMBB1000915, HEMBB1000975, HEMBB1001112, HEMBB1001151, HEMBB1001177, HEMBB1001302, HEMBB1001348, HEMBB1001564, HEMBB1001630, HEMBB1001871, HEMBB1001872, HEMBB1001925, HEMBB1001962, HEMBB1002042, HEMBB1002044, HEMBB1002142, HEMBB1002190, HEMBB1002193, HEMBB1002247. HEMBB1002383. HEMBB1002387. HEMBB1002550. HEMBB1002600. HEMBB1002692. MAMMA1000045, MAMMA1000129, MAMMA1000133, MAMMA1000277, MAMMA1000278, MAMMA1000410, MAMMA1000416. MAMMA1000472. MAMMA1000672. MAMMA1000684. MAMMA1000714. MAMMA1000734. MAMMA1000778, MAMMA1000798, MAMMA1000842, MAMMA1000859, MAMMA1000897, MAMMA1000956, MAMMA1001008, MAMMA1001030, MAMMA1001041, MAMMA1001073, MAMMA1001080, MAMMA1001139, MAMMA1001154. MAMMA1001322. MAMMA1001388. MAMMA1001411. MAMMA1001487. MAMMA1001751. MAMMA1001754, MAMMA1001771, MAMMA1002009, MAMMA1002427, MAMMA1002428, MAMMA1002461, MAMMA1002524, MAMMA1002573, MAMMA1002598, MAMMA1002655, MAMMA1002684, MAMMA1002769, MAMMA1002844. MAMMA1002881. MAMMA1002890. MAMMA1002938. MAMMA1002947. MAMMA1003035. MAMMA1003089, MAMMA1003146, MAMMA1003150, NT2RM1000035, NT2RM1000037, NT2RM1000062, NT2RM1000080, NT2RM1000092, NT2RM1000131, NT2RM1000199, NT2RM1000257, NT2RM1000260, NT2RM1000355. NT2RM1000430. NT2RM1000563. NT2RM1000648. NT2RM1000742. NT2RM1000770. NT2RM1000800. NT2RM1000811. NT2RM1000833. NT2RM1000857. NT2RM1000867. NT2RM1000882. NT2RM1000905, NT2RM1001008, NT2RM1001115. NT2RM1001139. NT2RM2000259. NT2RM2000260. NT2RM2000287. NT2RM2000395. NT2RM2000402, NT2RM2000407, NT2RM2000422, NT2RM2000490, NT2RM2000522, NT2RM2000566,

	NT2RM2000581,	NT2RM2000609,	NT2RM2000821,	NT2RM2001370,	NT2RM2001393,	NT2RM2001499,
	NT2RM2001547,	NT2RM2001613,	NT2RM2001648,	NT2RM2001659,	NT2RM2001671,	NT2RM2001688,
	NT2RM2001698,	NT2RM2001718,	NT2RM2001753,	NT2RM2001760,	NT2RM2001785,	NT2RM2001930,
	NT2RM2001950,	NT2RM2001997,	NT2RM2001998,	NT2RM2002049,	NT2RM2002145,	NT2RM4000233,
5	NT2RM4000433,	NT2RM4000457,	NT2RM4000486,	NT2RM4000496,	NT2RM4000520,	NT2RM4000634,
	NT2RM4000674,	NT2RM4000700,	NT2RM4000764,	NT2RM4000778,	NT2RM4000795,	NT2RM4000820,
	NT2RM4000857,	NT2RM4001032,	NT2RM4001054,	NT2RM4001116,	NT2RM4001455,	NT2RM4001666,
	NT2RM4001810,	NT2RM4001813,	NT2RM4001930,	NT2RM4001987,	NT2RM4002054,	NT2RM4002073,
	NT2RM4002145,	NT2RM4002146,	NT2RM4002189,	NT2RM4002194,	NT2RM4002251,	NT2RM4002339,
10	NT2RM4002438,	NT2RM4002446,	NT2RM4002452,	NT2RM4002460,	NT2RM4002493,	NT2RM4002558,
	NT2RM4002565.	NT2RM4002571.	NT2RM4002594,	NT2RP1000130.	NT2RP1000191.	NT2RP1000326,
	NT2RP1000358,	NT2RP1000413,	NT2RP1000418,	NT2RP1000547,	NT2RP1000609,	NT2RP1000677,
	NT2RP1000767,	NT2RP1000782,	NT2RP1000856,	NT2RP1001113,	NT2RP1001247,	NT2RP1001286,
	NT2RP1001310.	NT2RP1001311.	NT2RP1001313.	NT2RP1001385.	NT2RP1001449.	NT2RP1001546.
15	NT2RP1001569,	NT2RP2000032,	NT2RP2000040,	NT2RP2000056,	NT2RP2000070,	NT2RP2000091,
	NT2RP2000114	NT2RP2000120,	NT2RP2000173,	NT2RP2000175	NT2RP2000195,	NT2RP2000257,
	NT2RP2000270.	NT2RP2000283,	NT2RP2000288.	NT2RP2000289.	NT2RP2000459.	NT2RP2000516.
	NT2RP2000660,	NT2RP2000842,	NT2RP2000892,	NT2RP2001081,	NT2RP2001268,	NT2RP2001295,
	NT2RP2001366.	NT2RP2001378.	NT2RP2001576.	NT2RP2001581,	NT2RP2001597,	NT2RP2001613.
20	NT2RP2001947,	NT2RP2001991,	NT2RP2002025,	NT2RP2002066.	NT2RP2002078,	NT2RP2002105.
	NT2RP2002312.	NT2RP2002325.	NT2RP2002385.	NT2RP2002479.	NT2RP2002537.	NT2RP2002643.
	NT2RP2002701.	NT2RP2002740.	NT2RP2002857.	NT2RP2003125.	NT2RP2003297,	NT2RP2003433.
	NT2RP2003446.	NT2RP2003466,	NT2RP2003506,	NT2RP2003513.	NT2RP2003629.	NT2RP2003668.
	NT2RP2003760,	NT2RP2003777,	NT2RP2003781,	NT2RP2004041,	NT2RP2004142,	NT2RP2004194,
25	NT2RP2004270,	NT2RP2004300,	NT2RP2004392,	NT2RP2004655.	NT2RP2004681,	NT2RP2004775,
	NT2RP2004799,	NT2RP2004936,	NT2RP2004959,	NT2RP2005012,	NT2RP2005159,	NT2RP2005227,
	NT2RP2005270,	NT2RP2005344,	NT2RP2005465,	NT2RP2005509,	NT2RP2005752,	NT2RP2005781,
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	NT2RP2006261,	NT2RP2006565,	NT2RP2006571,	NT2RP2006573.	NT2RP3000092.	NT2RP3000109.
30	NT2RP3000134,	NT2RP3000207,	NT2RP3000333,	NT2RP3000341,	NT2RP3000393,	NT2RP3000439,
	NT2RP3000441.	NT2RP3000531.	NT2RP3000685.	NT2RP3000825.	NT2RP3000826.	NT2RP3000852.
	NT2RP3000919,	NT2RP3001084,	NT2RP3001096,	NT2RP3001126,	NT2RP3001140,	NT2RP3001176,
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	NT2RP3001497.	NT2RP3001538,	NT2RP3001589.	NT2RP3001642.	NT2RP3001708,	NT2RP3001716.
35	NT2RP3001727,	NT2RP3001739,	NT2RP3001799,	NT2RP3001943.	NT2RP3001944,	NT2RP3002002,
	NT2RP3002007,	NT2RP3002014,	NT2RP3002054,	NT2RP3002108,	NT2RP3002163,	NT2RP3002351,
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	NT2RP3003665,	NT2RP3003672,	NT2RP3003701,	NT2RP3003716.	NT2RP3003799,	NT2RP3003828.
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45	NT2RP4000417.	NT2RP4000500.	NT2RP4000524.	NT2RP4000556.	NT2RP4000560,	NT2RP4000588.
	NT2RP4000713,	NT2RP4000724,	NT2RP4000817,	NT2RP4000833,	NT2RP4000878,	NT2RP4000907,
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	NT2RP4001219.	NT2RP4001274.	NT2RP4001313.	NT2RP4001345.	NT2RP4001372.	NT2RP4001373.
50	NT2RP4001379,	NT2RP4001498,	NT2RP4001547,	NT2RP4001571,	NT2RP4001574,	NT2RP4001644,
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	NT2RP4001823.	NT2RP4001950.	NT2RP4001730, NT2RP4001975.	NT2RP4001739, NT2RP4002052.	NT2RP4001803, NT2RP4002075.	NT2RP5003500.
				RC1000060, OVAR		
	1000689. OVAR				VARC 1000850,	OVARC1000890.
55	OVARC1000924,	OVARC1000936,	OVARC1000959,	OVARC1000751, C	OVARC1000999,	OVARC1000690,
~	OVARC1000924,	OVARC1000936,	OVARC1000939,	OVARC1000364,	OVARC1000999,	OVARC1001034,
			OVARC1001129, OVARC1001476,			
	OVARC1001391, OVARC1001703,	OVARC1001453, OVARC1001713.	OVARC1001476, OVARC1001745.	OVARC1001506, OVARC1001767.	OVARC1001610, OVARC1002127,	OVARC1001702, OVARC1002138.
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                                                                              PLACE1005898.
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                                 PLACE1006678.
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                                                               PLACE2000477.
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                                                PLACE4000259.
                                                               PLACE4000300,
                                                                              PLACE4000387.
    PLACE4000431.
                   PLACE4000487.
                                 PLACE4000494.
                                                PLACE4000522.
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                                                               THYRO1000394.
    PLACE4000593.
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                                                THYRO1000327.
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                                 THYRO1000756.
                                                THYRO1000783.
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                                 THYRO1001401. THYRO1001534.
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    THYRO1001828.
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                                                Y79AA1001384
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                   Y79AA1001023
                                  Y79AA1001177,
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    Y79AA1001647.
                   Y79AA1001846.
                                 Y79AA1001874.
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                                                              Y79AA1002246.
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    Y79AA1002399, Y79AA1002416.
    [0276] The following 141 clones are categorized into glycoproteins-associated proteins.
    HEMBA1000156. HEMBA1000518. HEMBA1000852. HEMBA1001071. HEMBA1001286. HEMBA1001661.
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    HEMBA1003679, HEMBA1003866, HEMBA1005576, HEMBA1005581, HEMBA1005699, HEMBA1006038,
    HEMBA1006976, HEMBA1007301, HEMBB1000317, HEMBB1000915, HEMBB1001871, HEMBB1001872,
    HEMBB1002193, MAMMA1000672, MAMMA1000897, MAMMA1001030, MAMMA1001388, MAMMA1002329,
    MAMMA1002428, MAMMA1002573, MAMMA1003150, NT2RM1000648, NT2RM1001115, NT2RM2000260,
    NT2RM2000407.
                   NT2RM2000422, NT2RM2000490, NT2RM2001499, NT2RM2001659,
                                                                              NT2RM2001930.
    NT2RM4000820.
                   NT2RM4000857, NT2RM4001810, NT2RM4001813, NT2RM4001987,
                                                                              NT2RM4002145
                                 NT2RM4002460, NT2RM4002558, NT2RP1000677,
    NT2RM4002189, NT2RM4002251,
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    NT2RP1000856.
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    NT2RP2001597.
                   NT2RP2001991.
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    NT2RP2004732
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    NT2RP3002810.
                   NT2RP3003672.
                                 NT2RP3003701. NT2RP3003716. NT2RP3003914.
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    NT2RP4000212.
                   NT2RP4000417.
                                 NT2RP4000724. NT2RP4000817. NT2RP4000925.
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    NT2RP4001372.
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PLACE1006534, PLACE1006754, PLACE1006956, PLACE1007416, PLACE1007632, PLACE1007649,
PLACE1008643, PLACE1009094, PLACE1009992, PLACE1010231, PLACE1010662, PLACE1011371,
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PLACE3000339. PLACE400063. PLACE4000230. PLACE4000522. PLACE4000548. PLACE4000581.
THYRO1000327, THYRO1000756, THYRO1001287, Y79AA1001603, Y79AA1001874
[0277] The following 129 clones are categorized into signal transduction-associated proteins.
HEMBA1000303, HEMBA1000369, HEMBA1000608, HEMBA1000657, HEMBA1000919, HEMBA1001019,
HEMBA1001174, HEMBA1001822, HEMBA1001921, HEMBA1002139, HEMBA1002212, HEMBA1002341,
HEMBA1002417, HEMBA1002768, HEMBA1003250, HEMBA1003291, HEMBA1003645, HEMBA1004286,
HEMBA1005737. HEMBA1006130. HEMBA1006708. HEMBB1000083. HEMBB1000266. HEMBB1000632.
HEMBB1000781, HEMBB1000831, HEMBB1002193, MAMMA1000173, MAMMA1001038, MAMMA1001198,
MAMMA1002842, MAMMA1003057, NT2RM1000702, NT2RM1000772, NT2RM1001072, NT2RM2000030,
NT2RM2000469, NT2RM2000612, NT2RM2001221, NT2RM2001345, NT2RM2002128, NT2RM4000229,
NT2RM4000354, NT2RM4000611, NT2RM4000798, NT2RM4001411, NT2RM4001412, NT2RM4001629,
NT2RM4001758, NT2RM4002013, NT2RM4002527, NT2RP1000018, NT2RP1000701, NT2RP1001294,
NT2RP1001302. NT2RP2000668. NT2RP2001440. NT2RP2001560. NT2RP2002058. NT2RP2002193.
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NT2RP3001857, NT2RP3001938, NT2RP3002004, NT2RP3002785, NT2RP3002909, NT2RP3002988,
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NT2RP4001896, NT2RP4001927, NT2RP4002408, NT2RP5003477, OVARC1000013, OVARC1000437,
OVARC1000556, OVARC1000649, OVARC 1000945, OVARC1001200, OVARC1002182, PLACE1000977,
PLACE1001387, PLACE1002493, PLACE1002591, PLACE1003190, PLACE1003353, PLACE1004128,
PLACE1004302, PLACE1004937, PLACE1005243, PLACE1008000, PLACE1008244, PLACE1008650,
PLACE1009468, PLACE1009596, PLACE1009708, PLACE1009845, PLACE1010926, PLACE1011041,
PLACE2000164. PLACE2000371, PLACE3000145, PLACE3000350, THYRO1000072, THYRO1000748,
THYRO1001120, Y79AA1000328, Y79AA1002431
[0278] The following 309 clones are categorized into transcription -associated proteins.
HEMBA1000158, HEMBA1000201, HEMBA1000216, HEMBA1000555, HEMBA1000561, HEMBA1000851,
HEMBA1001077, HEMBA1001137, HEMBA1001405, HEMBA1001510, HEMBA1001635, HEMBA1001804,
HEMBA1001809, HEMBA1001819, HEMBA1001847, HEMBA1001869, HEMBA1002035, HEMBA1002092,
HEMBA1002177, HEMBA1002770, HEMBA1002935, HEMBA1003408, HEMBA1003545, HEMBA1003568,
HEMBA1003662. HEMBA1003684. HEMBA1003760. HEMBA1003953. HEMBA1004097. HEMBA1004321.
HEMBA1004353, HEMBA1004389, HEMBA1004479, HEMBA1004758, HEMBA1004973, HEMBA1005219,
HEMBA1005359, HEMBA1005513, HEMBA1005528, HEMBA1005548, HEMBA1005558, HEMBA1005931,
HEMBA1006158. HEMBA1006248. HEMBA1006278. HEMBA1006283. HEMBA1006347. HEMBA1006359.
HEMBA1006559, HEMBA1006941, HEMBB1000789, HEMBB1001011, HEMBB1001314, HEMBB1001482,
HEMBB1001673, HEMBB1001749, HEMBB1001839, HEMBB1001908, HEMBB1002134, HEMBB1002217,
HEMBB1002342. HEMBB1002607. MAMMA1000183. MAMMA1000388. MAMMA1001105. MAMMA1001222.
MAMMA1001260. MAMMA1001627. MAMMA1001633. MAMMA1001743. MAMMA1001820. MAMMA1001837.
MAMMA1002617, MAMMA1002650, MAMMA1002937, NT2RM1000055, NT2RM1000086, NT2RM1000746,
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NT2RM2000740, NT2RM2001035, NT2RM2001105, NT2RM2001575, NT2RM2001670, NT2RM2001716.
NT2RM2001771, NT2RM2002091, NT2RM4000024, NT2RM4000046, NT2RM4000104, NT2RM4000202,
NT2RM4000531, NT2RM4000595, NT2RM4000733, NT2RM4000734.
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NT2RP2001233, NT2RP2001756, NT2RP2001869, NT2RP2002046, NT2RP2002252, NT2RP2002270,
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NT2RP2003564, NT2RP2003714, NT2RP2004013, NT2RP2004066, NT2RP2004187,
                                                                         NT2RP2004920.
NT2RP2004961. NT2RP2005003. NT2RP2005139. NT2RP2005325. NT2RP2005496. NT2RP2005701.
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OVARC1001271, OVARC1001417, OVARC1001436, PLACE1000133, PLACE1000583, PLACE1000706,
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PLACE1002532, PLACE1002775, PLACE1002834, PLACE1003302, PLACE1003605, PLACE1003738,
PLACE1003885. PLACE1004471. PLACE1005584. PLACE1005803. PLACE1005966. PLACE1006167.
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PLACE4000261, PLACE4000450, PLACE4000489, THYRO1000085, THYRO1000121, THYRO1000242,
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              Y79AA1000033, Y79AA1000037,
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Y79AA1001299.
              Y79AA1001312. Y79AA1001391.
                                         Y79AA1001533. Y79AA1001613. Y79AA1001866.
Y79AA1002103, Y79AA1002229, Y79AA1002433, Y79AA1002472, Y79AA1002482,
[0279] The following 392 clones are categorized into disease-associated proteins.
HEMBA1000020, HEMBA1000216, HEMBA1000304, HEMBA1000561, HEMBA1000569, HEMBA1000910,
HEMBA1001043, HEMBA1001059, HEMBA1001071, HEMBA1001088, HEMBA1001569, HEMBA1001661,
HEMBA1001672, HEMBA1001819, HEMBA1001921, HEMBA1002267, HEMBA1002419, HEMBA1002469,
HEMBA1002547. HEMBA1002555. HEMBA1002810. HEMBA1002939. HEMBA1002997. HEMBA1003148.
HEMBA1003369, HEMBA1003417, HEMBA1003418, HEMBA1003433, HEMBA1003538, HEMBA1003555,
HEMBA1003568, HEMBA1003569, HEMBA1003581, HEMBA1004168, HEMBA1004202, HEMBA1004248,
HEMBA1004275, HEMBA1004321, HEMBA1004353, HEMBA1004356, HEMBA1004479, HEMBA1004509,
HEMBA1004669, HEMBA1005009, HEMBA1005338, HEMBA1005367, HEMBA1005423, HEMBA1005528,
HEMBA1005581, HEMBA1005621, HEMBA1005699, HEMBA1006507, HEMBA1006650, HEMBA1006652,
HEMBA1006737, HEMBA1006807, HEMBA1006877, HEMBA1007121, HEMBA1007243, HEMBB1000119,
HEMBB1000693, HEMBB1000927, HEMBB1000985, HEMBB1001068, HEMBB1001282, HEMBB1001339,
HEMBB1001482, HEMBB1001564, HEMBB1001802, HEMBB1001905, HEMBB1001908, HEMBB1002217,
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NT2RP1001665, NT2RP2000070, NT2RP2000147, NT2RP2000224, NT2RP2000248,
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NT2RP2000310, NT2RP2000414, NT2RP2000420, NT2RP2000523, NT2RP2000809,
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NT2RP2004732 NT2RP2004933 NT2RP2005003 NT2RP2005144 NT2RP2005239 NT2RP2005276
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	NT2RP2005288,	NT2RP2005315,	NT2RP2005325,	NT2RP2005336,	NT2RP2005358,	NT2RP2005407,
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	NT2RP2006071,	NT2RP2006219,	NT2RP2006312,	NT2RP2006456,	NT2RP3000050,	NT2RP3000068,
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	NT2RP3002953,	NT2RP3002988,	NT2RP3003078,	NT2RP3003251,	NT2RP3003282,	NT2RP3003313,
	NT2RP3003327,	NT2RP3003409,	NT2RP3003672,	NT2RP3003831,	NT2RP3004016,	NT2RP3004078,
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	NT2RP4000925,	NT2RP4001086,	NT2RP4001126,	NT2RP4001150,	NT2RP4001213,	NT2RP4001276,
	NT2RP4001407,	NT2RP4001433,	NT2RP4001483,	NT2RP4001575,	NT2RP4001760,	NT2RP4001861,
	NT2RP4002078,	NT2RP4002791,	OVARC1000014,	OVARC1000139,	OVARC1000520,	OVARC1000722,
15	OVARC1000771,	OVARC1000834,	OVARC1001051,	OVARC1001113,	OVARC1001244,	OVARC1001372,
	OVARC1001417,	OVARC1001496,	OVARC1001506,	OVARC1001577,	OVARC1001726,	OVARC1001766,
	OVARC1001809,	OVARC1002165,	PLACE1000133,	PLACE1000383,	PLACE1000420,	PLACE1000583,
	PLACE1000588,	PLACE1001171,	PLACE1001387,	PLACE1001602,	PLACE1002046,	PLACE1002140,
	PLACE1002437,	PLACE1002474,	PLACE1002685,	PLACE1002782,	PLACE1002834,	PLACE1002908,
20	PLACE1003045,	PLACE1003302,	PLACE1003353,	PLACE1003366,	PLACE1003493,	PLACE1003669,
	PLACE1003704,	PLACE1003903,	PLACE1003968,	PLACE1004183,	PLACE1004197,	PLACE1004277,
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35	[0280] Among th	em Swiss-Prot dat	ahase search and G	enRank or LiniGene	database search re	vealed that the fol-

5 [0280] Among them, Swiss-Prot database search and GenBank or UniGene database search revealed that the following 380 clones matched the data of genes or proteins with had been registered in the database of Online Mendellan Inheritance in Man (OMIM) [http://www.ncbi.rlm.nih.gov/Omim/), which is a database of human genes and diseases. (The corresponding OMIM mumbers are parenthetically indicate following the cione names.)

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	Y79AA1002482(603971),  [0281] The following 425 clones presumably belong to enzymes and/or metabolism-associated proteins.					protoino
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[0282] The following 217 clones presumably belong to a group of cDNAs encoding ATP- and/or GTP-binding proteins.
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NT2RP4001592. NT2RP4001634. NT2RP4001644. NT2RP4001656. NT2RP4001896. NT2RP4002047.
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OVARC1000771, OVARC1000800, OVARC1001068, OVARC1002138, PLACE1000040, PLACE1000588,
PLACE1001104, PLACE1001739, PLACE1002433, PLACE1002437, PLACE1002714, PLACE1003394,
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PLACE2000404, PLACE2000427, PLACE3000350, PLACE4000009, PLACE4000014, PLACE4000326,
SKNMC1000013, THYRO1000072, THYRO1001458, Y79AA1000833, Y79AA1000962, Y79AA1001394,
Y79AA1001875, Y79AA1001963, Y79AA1002209,
102831 The following 320 clones presumably belong to nuclear proteins.
HEMBA1000005, HEMBA1000158, HEMBA1000216, HEMBA1000561, HEMBA1000591, HEMBA1001088,
HEMBA1001137, HEMBA1001405, HEMBA1001510, HEMBA1001579, HEMBA1001809, HEMBA1001819,
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HEMBA1002569, HEMBA1002935, HEMBA1002951, HEMBA1002999, HEMBA1003408, HEMBA1003545,
HEMBA1003662, HEMBA1003684, HEMBA1003690, HEMBA1003760, HEMBA1004203, HEMBA1004321,
HEMBA1004353. HEMBA1004479. HEMBA1004973. HEMBA1005219. HEMBA1005359. HEMBA1005558.
HEMBA1005931, HEMBA1006278, HEMBA1006283, HEMBA1006359, HEMBA1006485, HEMBA1007087,
HEMBB1000226, HEMBB1000789, HEMBB1001011, HEMBB1001056, HEMBB1001242, HEMBB1001482,
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MAMMA1001222. MAMMA1001260. MAMMA1001633. MAMMA1001743. MAMMA1001837. MAMMA1002617.
MAMMA1002869, MAMMA1002937, MAMMA1003011, NT2RM1000086, NT2RM1000187, NT2RM1000666,
NT2RM1000885. NT2RM1000894. NT2RM1001059. NT2RM1001092. NT2RM2000013. NT2RM2000588.
NT2RM2000624, NT2RM2000735, NT2RM2000740, NT2RM2001105, NT2RM2001635, NT2RM2001670,
NT2RM2001771, NT2RM2001823, NT2RM2001936, NT2RM2001989, NT2RM2002004, NT2RM2002088,
NT2RM2002091, NT2RM4000024, NT2RM4000046, NT2RM4000104, NT2RM4000202, NT2RM4000215,
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NT2RP2005701, NT2RP2005767,
                            NT2RP2005776, NT2RP2005933, NT2RP2005942,
                                                                       NT2RP2006043.
NT2RP2006436, NT2RP3000031, NT2RP3000050, NT2RP3000397, NT2RP3000512,
                                                                       NT2RP3000527
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NT2RP3000590. NT2RP3000603, NT2RP3000632, NT2RP3000917, NT2RP3001057,
NT2RP3001120. NT2RP3001253. NT2RP3001338. NT2RP3001384. NT2RP3001398.
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                            NT2RP3002399, NT2RP3002876, NT2RP3003193,
                                                                       NT2RP3003212
NT2RP3003555. NT2RP3004016. NT2RP3004206. NT2RP3004424. NT2RP3004428.
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NT2RP4000518, NT2RP4000997, NT2RP4001148, NT2RP4001206, NT2RP4001213,
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OVARC1000241, OVARC1000326, OVARC1000556, OVARC1000846, OVARC1001038, OVARC1001180,
OVARC1001232, OVARC1001271, OVARC1001306, OVARC1001436, OVARC1002112, PLACE1000133,
PLACE1000184, PLACE1000406,
                            PLACE1000583, PLACE1000596, PLACE1000979, PLACE1001118,
PLACE1001383, PLACE1001632, PLACE1002171, PLACE1002433, PLACE1002438, PLACE1002532,
PLACE1002775, PLACE1002816, PLACE1002834, PLACE1003100, PLACE1003190, PLACE1003302,
PLACE1003519. PLACE1003521. PLACE1003605. PLACE1003704. PLACE1003738.
                                                                       PLACE1003885,
PLACE1003923, PLACE1004302,
                            PLACE1004471, PLACE1004564, PLACE1004814,
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PLACE1005287. PLACE1005876. PLACE1005966. PLACE1006167. PLACE1006438.
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PLACE1006829. PLACE1006878. PLACE1006917. PLACE1007014. PLACE1007547.
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PLACE1007688. PLACE1007969. PLACE1008044. PLACE1008132. PLACE1008603.
                                                                       PLACE1009099.
PLACE1009130. PLACE1009308. PLACE1009398. PLACE1010134. PLACE1010194.
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PLACE1010720. PLACE1010870.
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                                          PLACE1011433, PLACE1011664,
                                                                       PLACE2000014.
PLACE2000427, PLACE3000009, PLACE3000169, PLACE4000014, PLACE4000156,
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PLACE4000261. PLACE4000326. PLACE4000489. SKNMC1000011. THYRO1000085. THYRO1000242.
THYRO1000585, THYRO1001100, THYRO1001189, THYRO1001809, Y79AA1000037, Y79AA1000214,
Y79AA1000231,
             Y79AA1000589, Y79AA1000752, Y79AA1001391, Y79AA1001613,
                                                                      Y79AA1001705
Y79AA1001963, Y79AA1002431, Y79AA1002472, Y79AA1002482
[0284] The following 292 clones presumably belong to DNA- and/or RNA-binding proteins.
HEMBA1000158, HEMBA1000216, HEMBA1000561, HEMBA1000591, HEMBA1000851, HEMBA1001088,
HEMBA1001137. HEMBA1001405. HEMBA1001510. HEMBA1001804. HEMBA1001809. HEMBA1001819.
HEMBA1001847. HEMBA1001869. HEMBA1002177. HEMBA1002935. HEMBA1003408. HEMBA1003545.
HEMBA1003568, HEMBA1003591, HEMBA1003662, HEMBA1003684, HEMBA1003760, HEMBA1003783,
HEMBA1003805, HEMBA1003953, HEMBA1004321, HEMBA1004354, HEMBA1004389, HEMBA1004479,
HEMBA1004669. HEMBA1004847. HEMBA1004973. HEMBA1005202. HEMBA1005359. HEMBA1005931.
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HEMBA1006248, HEMBA1006278, HEMBA1006283, HEMBA1006359, HEMBA1006652, HEMBA1007087,
HEMBA1007194, HEMBB1000264, HEMBB1000789, HEMBB1001011, HEMBB1001482, HEMBB1001736,
HEMBB1001749. HEMBB1001839. HEMBB1002217. MAMMA1000183. MAMMA1000284. MAMMA1000731.
MAMMA1001105. MAMMA1001222. MAMMA1001260. MAMMA1001743. MAMMA1001837. MAMMA1002385.
MAMMA1002617. MAMMA1002869. MAMMA1002937. MAMMA1003011. NT2RM1000086. NT2RM1000539.
NT2RM1000555. NT2RM1000666. NT2RM1000691. NT2RM1000826. NT2RM1000885. NT2RM1001059.
              NT2RM2000371, NT2RM2000624, NT2RM2000735, NT2RM2001105, NT2RM2001424,
NT2RM1001092.
NT2RM2001575,
              NT2RM2001605, NT2RM2001670, NT2RM2001771, NT2RM2001823,
                                                                         NT2RM2001989.
NT2RM2002004.
              NT2RM2002014. NT2RM2002088. NT2RM2002091. NT2RM4000046. NT2RM4000104.
NT2RM4000167.
              NT2RM4000191. NT2RM4000202. NT2RM4000531. NT2RM4000595. NT2RM4000733.
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NT2RP2001127.
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NT2RP2002079.
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NT2RP2003329.
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NT2RP2006043.
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OVARC1001987, OVARC1002112, PLACE1000406, PLACE1000583, PLACE1000979, PLACE1001118,
PLACE1001632,
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PLACE1007547,
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                                                                         PLACE1009099.
PLACE1009246, PLACE1009398,
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PLACE1010702. PLACE1010870.
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PLACE3000009.
                                           PLACE4000156.
                                                         PLACE4000192.
              PLACE3000169
                             PLACE4000014.
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PLACE4000489, SKNMC1000091, THYRO1000085, THYRO1000242, THYRO1000501,
                                                                         THYRO1001100
THYRO1001189. THYRO1001809. Y79AA1000037. Y79AA1000349.
                                                           Y79AA1000752.
                                                                          Y79AA1001211.
Y79AA1001312, Y79AA1001391, Y79AA1001613, Y79AA1002103, Y79AA1002472, Y79AA1002482,
[0285] The following 66 clones presumably belong to the category of RNA synthesis-associated proteins.
HEMBA1000591, HEMBA1001579, HEMBA1003179, HEMBA1003591, HEMBA1006278, HEMBB1000226,
NT2RM1000187.
              NT2RM1000852, NT2RM2000624, NT2RM2001989, NT2RM2002100, NT2RM4000191,
                                                          NT2RP1000470.
NT2RM4001178.
              NT2RM4002093.
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                                                                         NT2RP1001080.
NT2RP2000153.
              NT2RP2002928.
                             NT2RP2003157.
                                            NT2RP2004568.
                                                          NT2RP2005126.
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NT2RP2005539,
              NT2RP2005605.
                             NT2RP2005776, NT2RP2005942, NT2RP2006043,
                                                                         NT2RP2006238,
NT2RP3000361, NT2RP3000397,
                             NT2RP3001671,
                                            NT2RP3004504, NT2RP4000078,
                                                                         NT2RP4000111,
NT2RP4000481.
              NT2RP4000518. NT2RP4000614. NT2RP4000929. NT2RP4001696.
                                                                         NT2RP4002058.
OVARC1001232. OVARC1001577. PLACE1000406. PLACE1000596. PLACE1000755.
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PLACE1003704, PLACE1003885, PLACE1004564, PLACE1004814, PLACE1004902,
PLACE1005646. PLACE1005876. PLACE1006196. PLACE1006626. PLACE1006878.
                                                                         PLACE1006917.
PLACE1009476, PLACE1009925, PLACE1010194, PLACE1011114, THYRO1000121, Y79AA1001963,
[0286] The following 183 clones presumably belong to protein synthesis-associated and/or protein transport-asso-
ciated proteins
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HEMBA1000012, HEMBA1000141, HEMBA1000592, HEMBA1003617, HEMBA1003773, HEMBA1004202,

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HEMBA1004276, HEMBA1004734, HEMBA1004847, HEMBA1004929, HEMBA1004930, HEMBA1005047,
HEMBA1005202, HEMBA1006031, HEMBA1006272, HEMBA1006474, HEMBA1006652, HEMBA1006914,
HEMBA1006973, HEMBA1007224, HEMBB1000915, HEMBB1001112, HEMBB1001137, HEMBB1001736,
HEMBB1001831, HEMBB1001915, MAMMA1000085, MAMMA1000734, MAMMA1001008, MAMMA1002170,
MAMMA1002219, MAMMA1002236, MAMMA1002619, NT2RM1000661, NT2RM1000833, NT2RM2000092,
NT2RM2000504. NT2RM2000577. NT2RM2000821. NT2RM2001201. NT2RM2001592. NT2RM2001613.
NT2RM2001648, NT2RM2001730, NT2RM2001760, NT2RM2002055, NT2RM4000155, NT2RM4000169,
NT2RM4000344, NT2RM4000356, NT2RM4000421, NT2RM4000712, NT2RM4001054, NT2RM4001203,
NT2RM4001382, NT2RM4001444, NT2RM4002062, NT2RM4002205, _ NT2RM4002623, NT2RP1000326,
NT2RP1000522, NT2RP1000547, NT2RP1000746, NT2RP1000947, NT2RP1001569, NT2RP2000147,
NT2RP2000710, NT2RP2000880, NT2RP2000943, NT2RP2001290, NT2RP2001392,
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NT2RP2001613, NT2RP2001660, NT2RP2001740, NT2RP2002124, NT2RP2002606, NT2RP2002862,
NT2RP2002959, NT2RP2002980, NT2RP2003137, NT2RP2003158, NT2RP2003391, NT2RP2003394,
NT2RP2003401, NT2RP2003433, NT2RP2003704, NT2RP2003713, NT2RP2003737, NT2RP2003760,
NT2RP2003981. NT2RP2004366. NT2RP2004389. NT2RP2004791. NT2RP2005012. NT2RP2005116.
NT2RP2005360, NT2RP2005763, NT2RP2005784, NT2RP3000366,
NT2RP3000759. NT2RP3000968, NT2RP3001113. NT2RP3001690. NT2RP3002045. NT2RP3002151.
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NT2RP4001313, NT2RP4001315, NT2RP4001574, NT2RP4001592, OVARC1000013, OVARC1000071,
OVARC1000085, OVARC1000465, OVARC1000564, OVARC1000771, OVARC1000862, OVARC1001171,
OVARC1001180, OVARC1001342, PLACE1000007, PLACE1000061, PLACE1000081, PLACE1000492,
PLACE1000863, PLACE1001092, PLACE1001748, PLACE1002090, PLACE1003174, PLACE1003915,
PLACE1004104, PLACE1004270, PLACE1004743, PLACE1005557, PLACE1005813, PLACE1006170,
PLACE1006488, PLACE1006829, PLACE1007706, PLACE1007729, PLACE1008273, PLACE1008402,
PLACE1008790, PLACE1008813, PLACE1009094, PLACE1009130, PLACE1009477, PLACE1009721,
PLACE1009845, PLACE1010074, PLACE1010547, PLACE1011109, PLACE1011229, PLACE1011477,
PLACE1012031, PLACE2000404, PLACE3000059, PLACE3000121, PLACE4000269, PLACE4000654,
SKNMC1000011, THYRO1000983, THYRO1001003, THYRO1001313, Y79AA1000560, Y79AA1000784,
Y79AA1000968, Y79AA1001493, Y79AA1001875, Y79AA1002027, Y79AA1002209,
[0287] The following 130 clones presumably belong to cytoskeletal-associated proteins.
HEMBA1000156, HEMBA1000168, HEMBA1000411, HEMBA1000588, HEMBA1001043, HEMBA1001651,
HEMBA1001661. HEMBA1002102. HEMBA1002161. HEMBA1002939. HEMBA1003235. HEMBA1003581.
HEMBA1004499, HEMBA1004534, HEMBA1004697, HEMBA1004929, HEMBA1004972, HEMBA1005582,
HEMBA1005595, HEMBA1006344, HEMBA1006737, HEMBB1001175, HEMBB1001282, HEMBB1001562,
HEMBB1001802, MAMMA1000824, MAMMA1001041, MAMMA1001576, MAMMA1001679, MAMMA1001735,
MAMMA1002297, MAMMA1002351, MAMMA1002622, MAMMA1002637, MAMMA1003127, NT2RM1000850,
NT2RM1000898, NT2RM2000030, NT2RM2000260, NT2RM2000691, NT2RM2001324, NT2RM4000169,
NT2RM4000229, NT2RM4000515, NT2RM4001217, NT2RP1000202, NT2RP1000348, NT2RP1000460,
NT2RP1000478, NT2RP1001033, NT2RP1001294, NT2RP1001302, NT2RP2000070, NT2RP2000812,
NT2RP2000814, NT2RP2001168, NT2RP2001245, NT2RP2001634, NT2RP2001900,
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NT2RP2003394. NT2RP2004041. NT2RP2004242. NT2RP2004538. NT2RP2004587.
                                                                        NT2RP2004681.
NT2RP2004732. NT2RP2004978. NT2RP2005491. NT2RP2005531. NT2RP2005712.
                                                                        NT2RP2006275.
NT2RP3000753. NT2RP3001113. NT2RP3001216. NT2RP3001239. NT2RP3001272.
                                                                        NT2RP3001554
NT2RP3001690. NT2RP3001799. NT2RP3002688. NT2RP3003061. NT2RP3003185.
                                                                        NT2RP3003230.
NT2RP3004569, NT2RP3004578, NT2RP4001004, NT2RP4001086, NT2RP4001256,
                                                                        NT2RP4001567.
NT2RP4001927, OVARC1000001, OVARC1000106, OVARC1000437, OVARC1000520, OVARC1000679,
OVARC1001731, OVARC1002050, PLACE1001104, PLACE1002571,
PLACE1002591, PLACE1002655, PLACE1002714, PLACE1003625, PLACE1005287, PLACE1006552,
PLACE1007946, PLACE1008426, PLACE1010148, PLACE1010547, PLACE1010743, PLACE1010896,
PLACE1010960, PLACE1011310, PLACE1011922, PLACE2000216, PLACE2000274, PLACE2000371,
PLACE2000458. PLACE3000145. PLACE3000416. PLACE4000009. THYRO1000132. THYRO1001405.
THYRO1001458, Y79AA1000368, Y79AA1000794, Y79AA1000833, Y79AA1000962, Y79AA1002208,
[0288] The following 54 clones presumably belong to cell division-associated and/or cell proliferation-associated
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HEMBA1001019, HEMBA1001595, HEMBA1002363, HEMBA1002997, HEMBA1003136, HEMBA1003369, HEMBA1004131, HEMBA1004504, HEMBA10045021, HEMBA10030037, HEMBB1000264, MAMMA1001768, M7ZRM1000348 NTZRM1000348 NTZRM1000364 NTZRM100031256 NTZRM2001743.

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NT2RM2001896, NT2RM2002145, NT2RM4000215, NT2RM4001714, NT2RP1000163, NT2RP1000333,
NT2RP1000439, NT2RP2000346, NT2RP2001397, NT2RP2002595, NT2RP2003177, NT2RP2003596,
NT2RP2003912, NT2RP2004396, NT2RP2005037, NT2RP2005520, NT2RP2005669,
                                                                                NT2RP2005835.
NT2RP3001730, NT2RP3002081, NT2RP4000210, NT2RP4000415, NT2RP4001414, NT2RP4001634,
OVARC1000013, OVARC1000937, PLACE1001383, PLACE1002433, PLACE1004316, PLACE1005287,
PLACE1008808, PLACE1010720, PLACE1010833, Y79AA1000748, Y79AA1001236, Y79AA1001394.
[0289] The following 36 clones presumably belong to the category of embryogenesis- and/or development-associ-
ated proteins
HEMBA1000518, HEMBA1001847, HEMBA1001869, HEMBA1003545, HEMBA1004973, HEMBB1002442,
MAMMA1001837, NT2RM2001670, NT2RM4000046, NT2RM4000531, NT2RM4001140, NT2RM4001858,
NT2RP2002078, NT2RP2004187, NT2RP2006436, NT2RP3000603, NT2RP3000994, NT2RP3001580,
NT2RP3001708, NT2RP3003071, NT2RP3004472, NT2RP3004617, NT2RP4000246, NT2RP4001567,
OVARC1000304, OVARC1000746, PLACE1000793, PLACE1002532, PLACE1003258, PLACE1003625,
PLACE1004460, PLACE1009622, PLACE4000558, THYRO1000085, Y79AA1001391, Y79AA1001692,
[0290] The following 30 clones presumably belong to cellular defense-associated proteins.
HEMBA1000005, HEMBA1000531, HEMBA1003417, HEMBA1006253, NT2RM4000354, NT2RM4001880,
NT2RP1000333, NT2RP1000493, NT2RP2000006, NT2RP2000045, NT2RP2000809, NT2RP2001536,
NT2RP2002464. NT2RP2004920. NT2RP2005037. NT2RP3000590. NT2RP3001426.
NT2RP3002785. NT2RP3004262. NT2RP4001555. NT2RP4001638. PLACE1006958.
                                                                                 PLACE1008275.
PLACE1009113, PLACE1011858, PLACE4000014, THYRO1000684, Y79AA1002139, Y79AA1002229,
[0291] Although it is unclear whether or not 261 clones out of clones other than the above-mentioned clones belong
to any of the above-described categories, these clones are predicted to have some functions, based on the homology
search using the full-length sequences thereof. The clone names and the gene definitions found in the result of ho-
mology search are shown below, separated with a double-slash mark, //.
   HEMBA1000030//Homo sapiens ARF GTPase-activating protein GIT1 mRNA, complete cds.
   HEMBA1000307//CARNITINE DEFICIENCY-ASSOCIATED PROTEIN EXPRESSED IN VENTRICLE 1
   HEMBA1000333//Homo sapiens F-box protein Fbx21 (FBX21) mRNA, complete cds.
   HEMBA1000488//RING CANAL PROTEIN (KELCH PROTEIN).
   HEMBA1000523//TESTIS-SPECIFIC PROTEIN PBS13.
   HEMBA1001197//Homo sapiens rap2 interacting protein x mRNA, complete cds.
   HEMBA1001302//Homo sapiens calcium binding protein precursor, mRNA, complete cds.
   HEMBA1001455//Mus musculus transposon-derived Buster2 transposase-like protein gene, partial cds.
   HEMBA1001675//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS9.
   HEMBA1001714//Homo sapiens mRNA for ATPase inhibitor precursor, complete cds.
   HEMBA1001744//SCY1 PROTEIN.
   HEMBA1001967//Homo sapiens NY-REN-57 antigen mRNA, partial cds.
   HEMBA1002151//Rattus norvegicus p34 mRNA, complete cds.
   HEMBA1002215//TESTIN 2 (TES2) [CONTAINS: TESTIN 1 (TES1)].
   HEMBA1002458//OVARIAN GRANULOSA CELL 13.0 KD PROTEIN HGR74.
   HEMBA1002777//Fugu rubripes BAW (BAW) mRNA, complete cds.
   HEMBA1003098//Homo sapiens NY-REN-6 antigen mRNA, partial cds.
   HEMBA1003199//Homo sapiens chromosome 5 F-box protein Fbx4 (FBX4) mRNA, complete cds.
   HEMBA1003615//Homo sapiens ART-4 mRNA, complete cds.
   HEMBA1003836//MOB1 PROTEIN (MPS1 BINDER 1).
   HEMBA1004295//Homo sapiens NY-REN-25 antigen mRNA, partial cds.
   HEMBA1004573//Homo sapiens mRNA for HELG protein.
   HEMBA1004604//Homo sapiens COP9 complex subunit 7a mRNA, complete cds.
   HEMBA1004795//CDC4-LIKE PROTEIN (FRAGMENT).
   HEMBA1005101//Homo sapiens SYT interacting protein SIP mRNA, complete cds.
   HEMBA1005201//Homo sapiens CGI-07 protein mRNA, complete cds.
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HEMBA1006268//Homo sapiens HQOO24c mRNA, complete cds.

HEMBA1005990//Homo sapiens I-1 receptor candidate protein mRNA, complete cds.

HEMBA1005206//Drosophila simulans anon73B1 gene and Su(P) gene.

HEMBA1005666//Homo sapiens mRNA for DIPB protein.

HEMBA1005530//Homo sapiens anaphase-promoting complex subunit 7 (APC7) mRNA, complete cds.

HEMBA1006398//Human L1 element L1.6 putative p150 gene, complete cds.

HEMBA1006445//Homo sapiens putative tumor supressor NOEY2 mRNA, complete cds.

HEMBA1007174//Homo sapiens epsin 2b mRNA, complete cds.

HEMBA1007251//Homo sapiens F-box protein FBX29 (FBX29) mRNA, partial cds. HEMBB1000036//Homo sapiens CGI-51 protein mRNA, complete cds.

HEMBB1000144//GUANYLATE CYCLASE ACTIVATING PROTEIN 2 (GCAP 2) (RETINAL

GUANYLYL CYCLASE ACTIVATOR PROTEIN P24).

HEMBB1000973//Mus musculus schlafen3 (Slfn3) mRNA, complete cds.

HEMBB1001058//Homo sapiens neuronal thread protein AD7c-NTP mRNA, complete cds

HEMBB1001234//65 KD YES-ASSOCIATED PROTEIN (YAP65).

HEMBB1001288//COPPER HOMEOSTASIS PROTEIN CUTC.

HEMBB1001331//Mus musculus mRNA for hepatoma-derived growth factor, complete cds, strain:BALB/c.

HEMBB1001384//Homo sapiens COP9 complex subunit 4 mRNA, complete cds.

HEMBR1002266//NEURONAL PROTEIN

15 HEMBB1002510//GYP7 PROTEIN.

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HEMBB1002705//Homo sapiens CGI-27 protein mRNA, complete cds.

MAMMA1000055//TESTIN 2 (TES2) [CONTAINS: TESTIN 1 (TEST)].

MAMMA1000625//GYP7 PROTEIN.

MAMMA1001075//Homo saniens CGI-72 protein mRNA, complete cds.

MAMMA1001181//ABC1 PROTEIN HOMOLOG PRECURSOR

MAMMA1001259//Mus musculus F-box protein FBX18 mRNA, partial cds.

MAMMA1001730//Homo sapiens brain and nasopharyngeal carcinoma susceptibility protein NSG-x mRNA, partial

MAMMA1002143//Homo sapiens Cdc42 effector protein 4 mRNA, complete cds.

25 MAMMA1002699//Rattus norvegicus EH domain binding protein Epsin mRNA, complete cds.

MAMMA1002972//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS27.

MAMMA1003113//Mus musculus COP9 complex subunit 7a (COPS7a) mRNA, complete cds.

NT2RM1000118//CALCINEURIN B SUBUNIT (PROTEIN PHOSPHATASE 2B REGULATORY SUBUNIT) (CAL-CINEURIN REGULATORY SUBUNIT)

30 NT2RM1000186//CALCINEURIN B SUBUNIT (PROTEIN PHOSPHATASE 2B REGULATORY SUBUNIT) (CAL-CINEURIN REGULATORY SUBUNIT).

NT2RM1000244//Homo sapiens TRAF4 associated factor 1 mRNA, partial cds.

NT2RM1000421//RIBONUCLEASE INHIBITOR.

NT2RM1000499//Caenorhabditis elegans mRNA for centaurin gamma 1A.

NT2RM1000623//RIBONUCLEASE INHIBITOR.

NT2RM1000883//Homo sapiens I-1 receptor candidate protein mRNA, complete cds.

NT2RM2000502//Rattus norvegicus W3O7 mRNA, complete cds.

NT2RM2000599//Homo sapiens F-box protein Lilina (LILINA) mRNA, complete cds. NT2RM2000718//Homo sapiens endocrine regulator mRNA, complete cds.

NT2RM2001065//Homo sapiens COP9 complex subunit 4 mRNA, complete cds.

NT2RM2001196//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).

NT2RM2001983//Homo sapiens RGS-GAIP interacting protein GIPC mRNA, complete cds.

NT2RM2002109//Homo sapiens glioma amplified on chromosome 1 protein (GAC1) mRNA, complete cds.

NT2RM2002142//GASTRULATION SPECIFIC PROTEIN G12.

45 NT2RM4000030//LAS1 PROTEIN.

NT2RM4000139//R.norvegicus trg mRNA.

NT2RM4000156//H. sapiens HPBRII-7 gene.

NT2RM4000386//Mus musculus ODZ3 (Odz3) mRNA, partial cds.

NT2RM4000590//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RM4001047//MO25 PROTEIN.

NT2RM4001155//ADRENAL MEDULLA 50 KD PROTEIN.

NT2RM4001256//Xenopus laevis putative Zic3 binding protein mRNA, complete cds.

NT2RM4001320//Homo sapiens mRNA for Neuroblastoma, complete cds. NT2RM4001340//UTR4 PROTEIN (UNKNOWN TRANSCRIPT 4 PROTEIN).

NT2RM4001347//Homo sapiens NY-REN-25 antigen mRNA, partial cds.

NT2RM4001371//Homo sapiens IDN3 mRNA, partial cds.

NT2RM4001582//Mus musculus COP9 complex subunit 7b (COPS7b) mRNA, complete cds.

NT2RM4001611//SIS2 PROTEIN (HALOTOLERANCE PROTEIN HAL3).

NT2RM4001731//Homo sapiens F-box protein Lilina (LILINA) mRNA, complete cds.

NT2RM4001969//R.norvegicus mRNA for IP63 protein.

NT2RM4002034//Homo sapiens hiwi mRNA, partial cds.

NT2RM4002075//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RM4002226//GTPASE ACTIVATING PROTEIN ROTUND.

NT2RP1000040//Mus musculus donson protein (Donson) mRNA, partial cds.

NT2RP1000363//R.norvegicus LL5 mRNA.

NT2RP1000481//Homo sapiens antigen NY-CO-3 (NY-CO-3) mRNA, partial cds.

NT2RP1000513//Human NifU-like protein (hNifU) mRNA, partial cds.

NT2RP1000733//Human mRNA for GSPT1-TK protein,complete cds.

NT2RP1000860//Homo sapiens KLO4P mRNA, complete cds.

NT2RP1000954//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RP1001011//Drosophila melanogaster putative 43 kDa protein (TH1) mRNA, complete cds

NT2RP1001395//Homo sapiens COP9 complex subunit 7a mRNA, complete cds,

15 NT2RP1001457//Homo sapiens partial mRNA for beta-transducin family protein (putative).

NT2RP1001494//MALE STERILITY PROTEIN 2.

NT2RP2000054//Homo sapiens putative ring zinc finger protein NY-REN-43 antigen mRNA, complete cds.

NT2RP2000067//Mus musculus ODZ3 (Odz3) mRNA, partial cds.

NT2RP2000133//Homo sapiens Leman coiled-coil protein (LCCP) mRNA, complete cds.

NT2RP2000157//MLO2 PROTEIN

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NT2RP2000965//Homo sapiens mRNA for fls353, complete cds.

NT2RP2001839//SCY1 PROTEIN.

NT2RP2001883//Homo sapiens CGI-01 protein mRNA, complete cds.

NT2RP2001976//Mus musculus calmodulin-binding protein SHA1 (Sha1) mRNA, complete cds.

NT2RP2001985//Homo sapiens high-risk human papilloma viruses E6 oncoproteins targeted protein E6TP1 alpha mRNA complete cds

NT2RP2002185//Homo saniens ubiquilin mRNA, complete cds.

NT2RP2002442//HESA PROTEIN

30 NT2RP2002727//Rattus norvegicus tulip 2 mRNA, complete cds.

NT2RP2002741//Homo sapiens mRNA for Neuroblastoma, complete cds.

NT2RP2002986//Homo sapiens mRNA for Kelch motif containing protein, complete cds.

NT2RP2003121//Mus musculus enhancer of polycomb (Epc1) mRNA, complete cds.

NT2RP2003265//Homo sapiens CGI-53 protein mRNA, complete cds.

35 NT2RP2003272//Homo sapiens ubiquilin mRNA, complete cds.

NT2RP2003857//MYOTROPHIN (V-1 PROTEIN) (GRANULE CELL DIFFERENTIATION PROTEIN).

NT2RP2003871//Homo sapiens transposon-derived Buster1 transposase-like protein gene, complete cds.

NT2RP2004425//Mus musculus axotrophin mRNA, complete cds.

NT2RP2004476//Homo sapiens cyclin L ania-6a mRNA, complete cds. NT2RP2004710//Mus musculus formin binding protein 30 mRNA, complete cds.

NT2RP2004816//H58 PROTEIN.

NT2RP2005441//Homo sapiens hypothalamus protein HT002 mRNA, complete cds.

NT2RP2005490//Mus musculus D3Mm3e (D3Mm3e) mRNA, complete cds.

NT2RP2005490//Mus musculus D3Mm3e (D3Mm3e) mRNA, complete cos NT2RP2005620//Homo sapiens epsin 2a mRNA, complete cds.

45 NT2RP2005654//CYSTEINE STRING PROTEIN (CCCS1).

NT2RP2005675//Homo sapiens growth suppressor related (DOC-1R) mRNA, complete cds.

NT2RP2005753//Homo sapiens I-1 receptor candidate protein mRNA, complete cds.

NT2RP2005841//Homo sapiens mRNA for ALEX3, complete cds.

NTZICE 200304 I/Fronto Sapiens HINNA for ALEXS, complete cus.

NT2RP2006598//Homo sapiens retinoid x receptor interacting protein mRNA, complete cds.

NT2RP3000047//NPL4 PROTEIN.

NT2RP3000233//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RP3000868//Human ovarian cancer downregulated myosin heavy chain homolog (Doc1) mRNA, complete

NT2RP3000869//Drosophila melanogaster AAA family protein Bor (bor) mRNA, complete cds.

55 NT2RP3001399//SSU72 PROTEIN.

NT2RP3001407//SCY1 PROTEIN.

NT2RP3001457//Drosophila melanogaster Melted (melt) mRNA, partial cds.

NT2RP3001587//Human anthracycline-associated resistance ARX mRNA, complete cds.

NT2RP3001712//Homo saniens HP1-BP74 protein mRNA, complete cds.

NT2RP3001819//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RP3001854//Homo sapiens novel retinal pigment epithelial cell protein (NORPEG) mRNA, complete cds.

NT2RP3001931//Rattus norvegicus clone C48 CDK5 activator-binding protein mRNA, complete cds.

NT2RP3002273//SCD6 PROTEIN

NT2RP3002631//Homo sapiens Ran binding protein 11 mRNA, complete cds.

NT2RP3002682//Homo saniens CGI-145 protein mRNA, complete cds.

NT2RP3002770//MYELOID DIFFERENTIATION PRIMARY RESPONSE PROTEIN MYD116

NT2RP3002818//INSERTION ELEMENT IS2A HYPOTHETICAL 48.2 KD PROTEIN.

NT2RP3002948//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RP3002972//Halocynthia roretzi mRNA for HrPET-1, complete cds,

NT2RP3003032//Homo sapiens okadaic acid-inducible and cAMP-regulated phosphoprotein 19 (ARPP-19) mR-NA. complete cds.

NT2RP3003290//Mus musculus mRNA for Ndr1 related protein Ndr3, complete cds.

15 NT2RP3003411//Mus musculus COP9 complex subunit 7b (COPS7b) mRNA, complete cds,

NT2RP3003491//Drosophila melanogaster Pelle associated protein Pellino (Pli) mRNA, complete cds.

NT2RP3003500//SCY1 PROTEIN.

NT2RP3003726//Homo sapiens spermatogenesis associated PD1 mRNA, complete cds.

NT2RP3004348//R. norvegicus mRNA for cytosolic resiniferatoxin-binding protein.

NT2RP3004507//MOB1 PROTEIN (MPS1 BINDER 1).

NT2RP4000129//Xenopus laevis F-box protein 28 (Fbx28) mRNA, partial cds.

NT2RP4000498//MOB1 PROTEIN (MPS1 BINDER 1).

NT2RP4000528//NPL4 PROTEIN

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NT2RP4000737//Mus musculus F-box protein FBL10 mRNA, partial cds.

25 NT2RP4000979//Homo sapiens putative HIV-1 infection related protein mRNA, partial cds.

NT2RP4001010//Rattus norvegicus PSD-95/SAP90-associated protein-4 mRNA, complete cds.

NT2RP4001207//Homo sapiens Ran binding protein 11 mRNA, complete cds.

NT2RP4001228//RING CANAL PROTEIN (KELCH PROTEIN).

NT2RP4001260//Homo sapiens F-box protein Fbx21 (FBX21) mRNA, complete cds.

NT2RP4001339//Homo sapiens mRNA for AMMERC1 protein.

NT2RP4001351//Human ovarian cancer downregulated myosin heavy chain homolog (Doc1) mRNA, complete

NT2RP4001474//Xenopus laevis putative Zic3 binding protein mRNA, complete cds.

NT2RP4001966//Mus musculus ODZ3 (Odz3) mRNA, partial cds. NT2RP4002018//RING CANAL PROTEIN (KELCH PROTEIN).

OVARC1000209//Oryza sativa submergence induced protein 2A mRNA, complete cds.

OVARC1000876//MOB1 PROTEIN (MPS1 BINDER 1).

OVARC1001065//Homo sapiens CGI-12 protein mRNA, complete cds.

OVARC1001092//Homo sapiens mRNA for JM5 protein, complete CDS (clone IMAGE 53337, LLNLc110F1857O7

40 (RZPD Berlin) and LLNLc110G0913Q7 (RZPD Berlin))

OVARC1001419//Homo sapiens GOK (STIM1) mRNA, complete cds.

OVARC1001555//NGG1-INTERACTING FACTOR 3.

OVARC1001711//CORNIFIN B (SMALL PROLINE-RICH PROTEIN 1B) (SPR1B) (SPR1 B).

OVARG1001943//Mus musculus DEBT-91 mRNA, complete cds. 45

PLACE1000004//Homo sapiens IDN3-B mRNA, complete cds.

PLACE1000066//SSU72 PROTEIN

PLACE1000610//MSN5 PROTEIN

PLACE1000636//MALE STERILITY PROTEIN 2.

PLACE1000769//Homo saniens CGI-18 protein mRNA, complete cds.

PLACE1000987//Rattus norvegicus late gestation lung 2 protein (Lgl2) mRNA, complete cds.

PLACE1001036//Homo sapiens mRNA for alpha integrin binding protein 63, partial.

PLACE1001845//Mus musculus cyclin ania-6a mRNA, complete cds.

PLACE1001920//Homo sapiens MDC-3,13 isoform 2 mRNA, complete cds.

PLACE1002665//Mus musculus enhancer of polycomb (Epc1) mRNA, complete cds.

PLACE1003602//Homo sapiens mRNA expressed in placenta.

PLACE1003611//Homo sapiens anaphase-promoting complex subunit 4 (APC4) mRNA, complete cds.

PLACE1004256//Mus musculus short coiled coil protein SCOCO (Scoc) mRNA, complete cds,

PLACE1004550//Homo sapiens CGI-20 protein mRNA, complete cds.

EP 1 074 617 A2 PLACE1004868//MALE STERILITY PROTEIN 2. PLACE1004930//Homo sapiens MDC-3.13 isoform 2 mRNA, complete cds PLACE1005052//Homo sapiens CGI-16 protein mRNA, complete cds. PLACE1005102//RING CANAL PROTEIN (KELCH PROTEIN). PLACE1005176//Homo sapiens hypothalamus protein HT001 mRNA, complete cds. PLACE1005187//APAG PROTEIN. PLACE1005331//Homo sapiens 7h3 protein mRNA, partial cds. PLACE1005727//Homo sapiens STRIN protein (STRIN) mRNA, complete cds. PLACE1006003//Homo sapiens CGI-94 protein mRNA, complete cds. PLACE1006335//Homo sapiens NY-REN-50 antigen mRNA, partial cds. PLACE1006385//Homo sapiens epsin 2a mRNA, complete cds. PLACE1006506//Homo sapiens anaphase-promoting complex subunit 4 (APC4) mRNA, complete cds. PLACE1007105//Homo sapiens muskelin (MKLN1) mRNA, complete cds. PLACE1007537//Homo sapiens ankyrin repeat-containing protein ASB-2 mRNA, complete cds. PLACE1007705//Mus musculus mRNA for Ndr1 related protein Ndr3, complete cds. PLACE1007791//Homo sapiens IDN3-B mRNA, complete cds. PLACE1007897//Homo sapiens FLASH mRNA, complete cds. PLACE1008080//Homo sapiens mRNA for HEXIM1 protein, complete cds. PLACE1008368//RING CANAL PROTEIN (KELCH PROTEIN). PLACE1008398//GENE 33 POLYPEPTIDE PLACE1008465//Homo sapiens mRNA for rapa-1 (rapa gene). PLACE1008627//Homo sapiens mRNA for cysteine-rich protein. PLACE 1009020 // NIES PROTEIN PLACE1009060//BRO1 PROTEIN PLACE1009186//Homo sapiens small zinc finger-like protein (TIM9b) mRNA, complete cds. PLACE1009443//Mus musculus F-box protein FBL8 mRNA, complete cds. PLACE1009571//Homo sapiens PTD002 mRNA, complete cds. PLACE1009670//Homo sapiens genethonin 1 mRNA, complete cds. PLACE1010105//RING CANAL PROTEIN (KELCH PROTEIN). PLACE1010261//SEGREGATION DISTORTER PROTEIN. PLACE1010310//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT). PLACE1010522//Homo sapiens mRNA for DEPP (decidual protein induced by progesterone), complete cds. PLACE1010579//Homo sapiens CED-6 protein (CED-6) mRNA, complete cds. PLACE1010628//Homo sapiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, partial cds. PLACE1010661//TESTIS-SPECIFIC PROTEIN PBS13. PLACE1010761//Homo sapiens mRNA for cisplatin resistance-associated overexpressed protein, complete cds. PLACE1011185//INSERTION ELEMENT IS1 PROTEIN INSB.

PLACE1011340//Homo sapiens IDN3-B mRNA, complete cds.

40 PLACE1011586//Rattus norvegicus clone C53 CDK5 activator-binding protein mRNA, complete cds.

PLACE2000246//RING CANAL PROTEIN (KELCH PROTEIN).

PLACE2000411//Homo sapiens epsin 2b mRNA, complete cds.

PLACE3000477//Homo sapiens phosphoprotein pp75 mRNA, partial cds.

THYRO1000173//Homo sapiens AP-mu chain family member mu1B (HSMU1B) mRNA, complete cds.

THYRO1000401//Human TcD37 homolog (HTcD37) mRNA, partial cds.

THYRO1000666//Mus musculus mRNA for kinesin like protein 9

THYRO1001033//TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521.

THYRO1001347//Homo sapiens RAN binding protein 16 mRNA, complete cds.

THYRO1001656//Homo sapiens Leman coiled-coil protein (LCCP) mRNA, complete cds.

THYRO1001703//NIFR3-LIKE PROTEIN

THYRO1001721//RING CANAL PROTEIN (KELCH PROTEIN).

Y79AA1000059//Homo sapiens aryl-hydrocarbon interacting protein-like 1 (AIPL1) gene, complete cds.

Y79AA1000181//Homo sapiens CGI-01 protein mRNA, complete cds.

Y79AA1000268//Mus musculus Nip2I mRNA, complete cds.

Y79AA1000313//CALPHOTIN.

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Y79AA1000540//CELL POLARITY PROTEIN TEA1

Y79AA1000966//Homo sapiens COP9 complex subunit 4 mRNA, complete cds.

Y79AA1000985//Human centrosomal protein kendrin mRNA, complete cds.

Y79AA1001323//Mus musculus mRNA for GSG1, complete cds.

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- Y79AA1001402//Homo sapiens paraneoplastic cancer-testis-brain antigen (MA4) mRNA, partial cds.
- Y79AA1001679//Homo sapiens lambda-crystallin mRNA, complete cds.
- Y79AA1001923//Homo sapiens F-box protein Fbx22 (FBX22) gene, partial cds. Y79AA1002083//H. sapiens mR-NA for MUF1 protein.
  - Y79AA1002307//Homo sapiens astrotactin2 (ASTN2) mRNA, complete cds.
  - Y79AA1002311//R. norvegicus mRNA for cytosolic resiniferatoxin-binding protein.
  - Y79AA1002487//Homo sapiens chromosome 5 F-box protein Fbx4 (FBX4) mRNA, complete cds.
- <sup>10</sup> [0292] Among the clones other than the above-mentioned, there were 36 clones that were similarly classified into the functional categories based on the results of functional domain search using the Pfam program. These clones were categorized as follows.
- [0293] Clones presumably belonging to the category of secretory or membrane proteins are two clones, MAMMA1002489 and NTZR-M002287; a clone presumably belonging to the category of glycoproteins-associated proteins are 11 clones, HeMBA100248: clones presumably belonging to the category of signal transduction-associated proteins are 11 clones, HeMBA1001247, NTZRP2001813, NTZRP200144, NTZRP2001940, NTZRP2003293, NTZRP20001487, NTZRP200311, PLACE1000127, e162100372, PLACE1003723, PLACE1005327, and PLACE30010124; ORDER 1001872, PLACE1001372, PLACE
- NTZRR4001235, PLACE1000560, PLACE1001716, PLACE1002499, and PLACE1007544; clones presumably belonging to the category of enzymes and/or metabolism-associated proteins are 2 clones, HEMBA1005732 and MAMMA1000402; clones presumably belonging to the category of DNA- and/or RNA-binding proteins are 4 clones, HEMBA1004596, OWARC1000144, PLACE1003334, and THYRO1001661; a clone presumably belonging to the category of protein synthesis- and/or protein transport-associated proteins is a clone, HEMBA1006294.
- 25 [0294] So far, useful information for presuming the functions is unavailable for the remaining 2511 clones. Their functions will possibly be revealed by further analyses. Names of the clones are listed below.

[0295] So far, useful information for presuming the functions are unavailable for the remaining 2511 clones. Their functions will possibly be revealed by further analyses. Names of the clones are listed below.

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	Y79AA1002220,	Y79AA1002234,	Y79AA1002298,	Y79AA1002407,	MAMMA1002215,	MAMMA1002721,

45 Homology Search Result Data 1.

[0296] The result of the homology search of the SwissProt using the 5'-end sequence.

[0297] Data include

NT2RP2002070.

the name of clone,

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definition of the top hit data,

the P-value: the length of the compared sequence: identity (%), and

the organism and the Accession No. of the top hit data, as in the order separated by //.

[0298] Data are not shown for the clones in which the P-value was higher than 1.

[0299] The P-value is a score obtained statistically by taking into account the possible similarity between two sequences. In general, the smaller P-value reflects the higher similarity, (Alischul, S.F., Gish, W., Miller, W., Myers, E. W., Samp; Lipman, D.J. (1990) Aquot;Basic local alignment search tool. Aquot; J. Mol. Biol. 215403-410; Gish, W. Samp;

States, D.J. (1993) "Identification of protein coding regions by database similarity search." Nature Genet. 3:266-272).

- F-HEMBA1000005//DNAJ PROTEIN HOMOLOG MTJ1/J1.8-85:244:75//MUS MUSCULUS (MOUSE)//O61712
  5 F-HEMBA1000012//PROBABLE LEUCYL-TRNA SYNTHETASE (EC 6.1.1.4) (LEUCINETRNA LIGASE) (LEURS)/J7.6e-57:231:53//CAENORHABDITIS ELEGANS //O09996
  - F-HEMBA1000020//TUBULIN BETA CHAIN.//1.0e-92:143:80//AJELLOMYCES CAPSULATA (HISTOPLASMA CAPSULATUM).//P41742
- F-HEMBA1000030//CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS).//0.021:136:33//PLASMODIUM KNOWLESI (STRAIN NURI).//P04922
  - F-HEMBA1000042//METALLOTHIONEIN 10-II (MT-10-II),//0.71:64:32//MYTILUS EDULIS (BLUE MUSSEL),// PR0247
  - F-HEMBA1000046//PROTEIN Q300.//0.92:40:37//MUS MUSCULUS (MOUSE).//Q02722
- F-HEMBA1000050//COMPETENCE PROTEIN S.//0.50:28:35//BACILLUS SUBTILIS.//P80355

  F-HEMBA1000076//ATP SYNTHASE E CHAIN. MITOCHONDRIAL (EC 3.6.1.34).//0.86:41:41//HOMO SAPIENS
- 15 F-HEMBA1000076/ATP SYNTHASE E CHAIN, MITOCHONDRIAL (EC 3.6.1.34).//0.86:41:41/HOMO SAPIENS (HUMAN)./P56385 F-HEMBA1000111
  - F-HEMBA1000129//UVSW PROTEIN (DAR PROTEIN).//0.023:68:33//BACTERIOPHAGE T4.//P20703
  - F-HEMBA1000141//YSY6 PROTEIN.//0.90:29:37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P38374
  - F-HEMBA1000150//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//8.4e-16:47:70//HOMO SAPIENS (HUMAN).//
    - F-HEMBA1000156//IMMEDIATE-EARLY PROTEIN.//8.1e-07:143:28//HERPESVIRUS SAIMIRI (STRAIN 11).// Q01042
- 25 F-HEMBA1000158//HYPOTHETICAL PROTEIN KIAA0192 (FRAGMENT),//7.9e-11:129:40//HOMO SAPIENS (HUMAN),//Q93074
  - F-HEMBA1000168//INSULIN RECEPTOR SUBSTRATE-2 (IRS-2) (4PS).//0.00055:86:36//MUS MUSCULUS (MOUSE).//P81122
- F-HEMBA1000180//VPU PROTEIN (U ORF PROTEIN).//0.22:73:28//CHIMPANZEE IMMUNODEFICIENCY VI-RUS (SIV(CPZ)) (CIV).//P17286
  - F-HEMBA1000185//RAS-1 PROTEIN.//5.1e-10:121:29//NEUROSPORA CRASSA.//P22126
    - F-HEMBA1000193//PROLINE-RICH PEPTIDE P-B.//0.00078:56:41//HOMO SAPIENS (HUMAN).//P02814
      F-HEMBA1000201//PROLINE-RICH PROTEIN MP-2 PRECURSOR //0.00061:49:42//MUS MUSCULUS
      (MOUSE).//P05142
  - F-HEMBA1000213
    - F-HEMBA1000216//HYPOXIA-INDUCIBLE FACTOR 1 ALPHA (HIF-1 ALPHA) (ARNT INTERACTING PROTEIN).//1.6e-59:115:53//MUS MUSCULUS (MOUSE).//Q61221
    - F-HEMBA1000227//SUPPRESSOR PROTEIN SRP40.//0.00059:135:22//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32583
- 40 F-HEMBA1000231//HYPOTHETICAL 60.7 KD PROTEIN C56F8.17C IN CHROMOSOME I.//0.024:60:38// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10264
  - F-HEMBA1000243//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.0038:125:34//HOMO SAPIENS (HU-MAN).//P08547
- F-HEMBA1000244//HYPOTHETICAL 123.6 KD PROTEIN IN POR2-COX5B INTERGENIC REGION.//3.1e-17: 149:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40480
  - F-HEMBA1000251
  - F-HEMBA1000264//PROBABLE E5 PROTEIN.//1.0:49:36//HUMAN PAPILLOMAVIRUS TYPE 58.//P26552
  - F-HEMBA1000280//SHORT NEUROTOXIN 1 (TOXIN C-6).//0.98:58:31//NAJA NAJA KAOUTHIA (MONOCLED COBRA) (NAJA NAJA SIAMENSIS).//P14613
  - F-HEMBA1000282//!!!! ALU SUBFAMILY J WARNING ENTRY!!!!/0.14:26:65//HOMO SAPIENS (HUMAN).//
    - F-HEMBA1000288 F-HEMBA1000290 SAPIENS (HUMAN F-HEMBA1000302
    - F-HEMBA1000290//HYPOTHETICAL 14 KD PROTEIN IN TVRI-6 REPETITIVE REGION.//3.8e-06:98:39//HOMO SAPIENS (HUMAN).//P10516
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- F-HEMBA1000303//HYPOTHETICAL 104.4 KD PROTEIN F54G8.4 IN CHROMOSOME III.//1.3e-05:69:42//
- F-HEMBA1000304//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/0.021:18:83//HOMO SAPIENS (HUMAN).//

## P39194

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F-HEMBA1000307//MYOSIN HEAVY CHAIN, CLONE 203 (FRAGMENT).//7.1e-06:235:25//HYDRA ATTENUATA (HYDRA) (HYDRA VULGARIS).//P39922

F-HEMBA1000327

F-HEMBA1000333//SRP1 PROTEIN.//1.0:159:30//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).// Q10193

F-HEMBA1000338//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//8.8e-26:36:83//HOMO SAPIENS (HUMAN).// P39193

E-HEMBA1000351

F-HEMBA1000355//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.99:22:50//HOMO SAPIENS (HUMAN).// P00841

F-HEMBA1000356//IMMEDIATE-EARLY PROTEIN IE180.//0.11:82:36//PSEUDORABIES VIRUS (STRAIN INDI-ANA-FUNKHAUSER / BECKER) (PRV).//P11675

F-HEMBA1000357///!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//2.1e-35:105:74//HOMO SAPIENS (HU-MAN).//P39192

F-HEMBA1000366//HYPOTHETICAL TRANSCRIPTIONAL REGULATOR AF1627.//1.0:28:42//ARCHAEOGLO-BUS FUL GIDUS //028646

F-HEMBA1000369//PRESYNAPTIC DENSITY PROTEIN 95 (PSD-95).//0.013:140:26//HOMO SAPIENS (HU-

20 F-HEMBA1000376//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49): ENDONUCLEASEI.//6.8e-08:66:42//MUS MUSCULUS (MOUSE)./P11369

F-HEMBA1000387//HYPOTHETICAL 63.2 KD PROTEIN C1F3.09 IN CHROMOSOME I.//1.5e-15:177:32// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10414

F-HEMBA1000390//PARATHYMOSIN.//0.0071:61:29//HOMO SAPIENS (HUMAN).//P20962

25 F-HEMBA1000392/!!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/2.1e-30:92:69//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1000396/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.9e-23:64:57//HOMO SAPIENS (HUMAN).//P08547

F-HEMBA1000411

30 F-HEMBA1000418

F-HEMBA1000422//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//8.3e-10:90:53//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1000428//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.1e-12:72:55//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1000434

F-HEMBA1000442//GENE 11 PROTEIN.//1.0:28:46//SPIROPLASMA VIRUS SPV1-R8A2 B.//P15902

F-HEMBA1000460//LYSIS PROTEIN (E PROTEIN) (GPE).//1.0:24:50//BACTERIOPHAGE ALPHA-3.//P31280

F-HEMBA1000469//PILI PROTEIN.//1.0:27:44//PSEUDOMONAS AERUGINOSA.//P43502

F-HEMBA1000488//ZINC FINGER PROTEIN 151 (MIZ-1 PROTEIN).//1.1e-07:90:38//HOMO SAPIENS (HU-MAN).//Q13105

F-HEMBA1000490//PLECTIN.//0.74:254:25//RATTUS NORVEGICUS (RAT).//P30427

F-HEMBA1000491//RAS-RELATED PROTEIN M-RAS.//3.0e-14:100:36//RATTUS NORVEGICUS (RAT).// P97538

F-HEMBA1000501/IIIII ALU SUBFAMILY SQ WARNING ENTRY IIIII/1.5e-20:81:54//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1000504

F-HEMBA1000505//NEURON-SPECIFIC X11 PROTEIN (FRAGMENT).//0.00028:128:32//HOMO SAPIENS (HUMAN).//Q02410

F-HEMBA1000508//CHITIN SYNTHASE 3 (EC 2.4.1.16) (CHITIN-UDP ACETYL-GLUCOSAMINYL TRANS-FERASE 3) //0.61:132:25//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P29465

F-HEMBA1000518

F-HEMBA1000519///!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.8e-37:68:75//HOMO SAPIENS (HUMAN).// P39189

- F-HEMBA1000520//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!/5.2e-09:75:49//HOMO SAPIENS (HUMAN).// P39192
- F-HEMBA1000523//TESTIS-SPECIFIC PROTEIN PBS13.//1.5e-35:257:36//MUS MUSCULUS (MOUSE).// On1755
- F-HEMBA1000531//HEAT SHOCK PROTEIN 70 B2.//1.6e-14:72:44//ANOPHELES ALBIMANUS (NEW WORLD MALARIA MOSQUITO).//P41827
  - F-HEMBA1000534//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//9.7e-32:96:78//HOMO SAPIENS (HUMAN).//
  - F-HEMBA1000540//LANTIBIOTIC LACTICIN 481 PRECURSOR (LACTOCOCCIN DR).//1.0:12:75//LACTOCOC-
    - CUS LACTIS (SUBSP. LACTIS) (STREPTOCOCCUS LACTIS)//P36499
      F-HEMBA1000542//SPERM MITOCHONDRIAL CAPSULE SELENOPROTEIN (MCS).//0.0089:79:31//MUS MUSCULUS (MOUSE)//P15265
  - F-HEMBA1000545//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//9.0e-83:256:66//HOMO SAPIENS (HU-MAN) //P08547
- 15 F-HEMBA1000555/JTRANSLATION INITIATION FACTOR IF-2.//3.6e-06:252:22//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P39730
  - F-HEMBA1000557
    - F-HEMBA1000561//ZINC FINGER PROTEIN 81 (FRAGMENT).//9.1 e-18:200:28//HOMO SAPIENS (HUMAN).// P51508
  - F-HEMBA1000563

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- F-HEMBA1000568
  - F-HEMBA1000569//GPI-ANCHORED PROTEIN P137.//1.0e-40:137:54//HOMO SAPIENS (HUMAN).//Q14444
  - F-HEMBA1000575
- F-HEMBA1000588
- 25 F-HEMBA1000591//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.1e-17:41:92//HOMO SAPIENS (HU-MAN) //P39194
  - F-HEMBA1000592//CENTROMERIC PROTEIN E (CENP-E PROTEIN).//0.18:128:23//HOMO SAPIENS (HU-MAN).//Q02224
- F-HEMBA1000594/HYPOTHETICAL 29.3 KD PROTEIN B0280.6 IN CHROMOSOME III.//0.93:24:54//
  CAENORHABDITIS ELEGANS.//P41997
  - F-HEMBA1000604//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00010:49:55//HOMO SAPIENS (HUMAN).// P39188
  - F-HEMBA1000608//HYPOTHETICAL PROTEIN KIAA0411 (FRAGMENT).//1.8e-55:179:61//HOMO SAPIENS (HUMAN).//O43295
- F-HEMBA1000622//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-21:94:62//HOMO SAPIENS (HUMAN).// P39188
  - F-HEMBA1000636//SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185).//0.34:73:36//VOLVOX CARTERI.// P21997
- F-HEMBA1000637//BASIC PROLINE-RICH PEPTIDE IB-1.//0.0057;76;38//HOMO SAPIENS (HUMAN).//P04281

  F-HEMBA1000655
  - F-HEMBA1000657//ZINC FINGER PROTEIN GCS1.//1.5e-07:66:37//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P35197
  - F-HEMBA1000662//METALLOTHIONEIN-II (MT-II).//0.79:33:39//CRICETULUS GRISEUS (CHINESE HAM-STER) //P02799
- 45 F-HEMBA1000673//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//3.1e-17:86:59//HOMO SAPIENS (HUMAN).// P39193
  - F-HEMBA1000682//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE].//3.0e-13:45:44//MUS MUSCULUS (MOUSE).//P11369
- F-HEMBA1000686/HYPOTHETICAL 48.0 KD PROTEIN C1B3.08 IN CHROMOSOME I.//4.5e-07:79:34//
  90 SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O13873
  - F-HEMBA1000702
    - F-HEMBA1000705//PROTEIN Q300.//0.80:25:44//MUS MUSCULUS (MOUSE).//Q02722
    - F-HEMBA1000719//MYOSIN IC HEAVY CHAIN.//0.0026:115:44//ACANTHAMOEBA CASTELLANII (AMOEBA).// P10569
- 55 F-HEMBA1000722
  - F-HEMBA1000726//!!!! ALU SUBFAMILY SB2 WARNING ENTRY!!!!//7.4e-32:83:77//HOMO SAPIENS (HU-MAN)///P39191
  - F-HEMBA1000727//ZINC FINGER PROTEIN CTH2 (YTIS11 PROTEIN),//0.73:26:46//SACCHAROMYCES CER-

EVISIAE (BAKER'S YEAST) //P47977

F-HEMBA1000747

F-HEMBA1000749//HYPOTHETICAL PROTEIN HI1484.//1.0:42:35//HAEMOPHILUS INFLUENZAE.//P44211 F-HEMBA1000752//RETROVIRUS-RELATED ENV POLYPROTEIN.//1.0e-08:84:39//HOMO SAPIENS (HU-

MAN) //P10267

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F-HEMBA 1000769

F-HEMBA1000773//PAIRED BOX PROTEIN PAX-4 //1.0:107:33//HOMO SAPIENS (HUMAN) //O43316 F-HEMBA1000774//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.3e-23:92:63//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1000791

F-HEMBA1000817//PROLACTIN RECEPTOR PRECURSOR (PRL-R),//0.079:87:29//CERVUS ELAPHUS (RED DEER),//Q28235

F-HEMBA1000822

F-HEMBA1000827//HYPOTHETICAL 8.4 KD PROTEIN.//0.98:48:39//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20546

15 F-HEMBA1000843//HYPOTHETICAL 7.3 KD PROTEIN D1044.5 IN CHROMOSOME III.//0.92:46:34// CAENORHARDITIS ELEGANS //P41953

F-HEMBA1000851//HOMEOBOX PROTEIN GBX-2 (GASTRULATION AND BRAIN-SPECIFIC HOMEOBOX PROTEIN 2).//0.048:39:51//HOMO SAPIENS (HUMAN).//P52951

F-HEMBA1000852//ARYLSULFATASE D PRECURSOR (EC 3.1.6.-) (ASD).//4.0e-24:29:100//HOMO SAPIENS (HUMAN).//P51689

F-HEMBA1000867

F-HEMBA1000869//PROBABLE E5 PROTEIN.//0.99;70;27//HUMAN PAPILLOMAVIRUS TYPE 18.//P06792 F-HEMBA1000870//MYOTOXIN 3 PRECURSOR (CROTAMINE 3),//0.79:43:32//CROTALUS DURISSUS TER-

RIFICUS (SOUTH AMERICAN RATTLESNAKE), //P24333 F-HEMBA1000872//GAR2 PROTEIN.//0.89:70:31//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST),//

P41891

F-HEMBA1000876//DEFENSIN.//0.89:34:38//ALLOMYRINA DICHOTOMA.//Q10745

F-HEMBA1000908//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS), //0.69:43:37//HO-30 MO SAPIENS (HUMAN),//P30808

F-HEMBA1000910//MELANOMA-ASSOCIATED ANTIGEN B3 (MAGE-B3 ANTIGEN).//5.1e-08:44:38//HOMO SAPIENS (HUMAN).//O15480

F-HEMBA1000918//60S RIBOSOMAL PROTEIN L37-A (YL35) (FRAGMENT).//1.0:19:52//SCHIZOSACCHARO-MYCES POMBE (FISSION YEAST), //P22667

F-HEMBA1000919//69 KD PARAFLAGELLAR ROD PROTEIN (69 KD PFR PROTEIN) (PFR-A/PFR-B).//0.29: 116:30//TRYPANOSOMA BRUCEI BRUCEI.//P22225 F-HEMBA1000934

F-HEMBA1000942/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/0.85:27:59//HOMO SAPIENS (HUMAN).// P39188

40 F-HEMBA1000943

F-HEMBA1000946//STO-2 PROTEIN //0.82:82:30//CAENORHABDITIS FLEGANS //Q19958

F-HEMBA1000960//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//0.0097;29;72//HOMO SAPIENS (HUMAN).// P30102

F-HEMBA1000968//METALLOTHIONEIN 20-III ISOFORMS A AND B (MT-20-IIIA AND MT-20-IIIB).//0.047:45:37// 45 MYTILUS EDULIS (BLUE MUSSEL).//P80253

F-HEMBA1000971//HYPOTHETICAL BHLF1 PROTEIN.//0.038:172:31//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181

F-HEMBA1000972

F-HEMBA1000974//HYPOTHETICAL PROTEIN MG441.//0.98:66:28//MYCOPLASMA GENITALIUM.//P47679 F-HEMBA1000975//COLLAGEN ALPHA 2(VIII) CHAIN (ENDOTHELIAL COLLAGEN) (FRAGMENT).//0.028:57: 36//HOMO SAPIENS (HUMAN),//P25067

F-HEMRA 1000985

F-HEMBA1000986//SUBMANDIBULAR GLAND SECRETORY GLX-RICH PROTEIN CB PRECURSOR (GRP-CB) (CONTIGUOUS REPEAT POLYPEPTIDE) (CRP),//0.13:91:34//RATTUS NORVEGICUS (RAT),//P08462

55 F-HEMBA1000991//HYPOTHETICAL 46.2 KD TRP-ASP REPEATS CONTAINING PROTEIN D2013.2 IN CHRO-MOSOME II.//5.6e-05:37:45//CAENORHABDITIS ELEGANS.//Q18964

F-HEMBA1001007//HYPOTHETICAL PROTEIN KIAA0179.//0.27:72:41//HOMO SAPIENS (HUMAN).//Q14684 F-HEMBA1001008//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/2.4e-25:61:70//HOMO SAPIENS (HU-

## MAN).//P39194

F-HEMBA1001009//CUTICLE COLLAGEN 34.//0.044:214:29//CAENORHABDITIS ELEGANS.//P34687

F-HEMBA1001017//SYNDECAN-3 PRECURSOR (N-SYNDECAN) (NEUROGLYCAN).//5.0e-85:191:84//RAT-TUS NORVEGICUS (RAT).//P33671

F-HEMBA1001019

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F-HEMBA1001020//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//6.7e-24:49:73//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1001022

F-HEMBA1001024//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.0e-11:61:59//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1001026/HYPOTHETICAL PROTEIN BB0073.//0.94:63:34//BORRELIA BURGDORFERI (LYME DIS-EASE SPIROCHETE).//O51100

F-HEMBA1001043//INVOLUCRIN.//0.0036:238:25//SAGUINUS OEDIPUS (COTTON-TOP TAMARIN),//P24712
F-HEMBA1001051//IIII ALU SUBFAMILY SB WARNING ENTRY !!!!//3.3e-32:95:75//HOMO SAPIENS (HUMAN).//

P39189
F-HEMBA1001052//CURROMYCIN RESISTANCE PROTEIN.//1.0:31:38//STREPTOMYCES HYGROSCOPI-

CUS.//P16961
F-HEMBA1001059/N-ACETYLGALACTOSAMINE-6-SULFATASE PRECURSOR (EC 3.1.6.4) (N- ACETYLGA-

LACTOSAMINE-6-SULFATS SULFATSS (CALACTOSE-6-SULFATSE) (CALACTOSAMINE-6-SULFATASE) (CALACTOSAMINE-6-SULFATASE) (CHONDROITINSULFATASE) (CHO

F-HEMBA1001071//PROCOLLAGEN ALPHA 1(III) CHAIN PRECURSOR.//8.3e-23:51:96//HOMO SAPIENS (HU-MAN) //P0/2461

F-HEMBA1001077//AUTOIMMUNE REGULATOR (APECED PROTEIN).//3.4e-06:37:56//HOMO SAPIENS (HU-MAN) //043018

F-HEMBA1001080/INFECTED CELL PROTEIN ICP34.5 (NEUROVIRULENCE FACTOR ICP34.5).//0.0012:70: 38//HERPES SIMPLEX VIRUS (TYPE 1 / STRAW MGH-10).//P37319

F-HEMBA1001085//SERINE/THREONINE PROTEIN PHOSPHATASE 5 (EC 3.1.3.16) (PPS) (PROTEIN PHOS-PHATASE T) (PPT) (FRAGMENT).//0.00018:76:32//MUS MUSCULUS (MOUSE).//Q60676

F-HEMBA1001088//PINCH PROTEIN (PARTICULARY INTERESTING NEW CYS-HIS PROTEIN).//3.5e-50:176: 57//HOMO SAPIENS (HUMAN).//P48059

F-HEMBA1001094

F-HEMBA1001099//LIGHT-HARVESTING PROTEIN B800/850/890, ALPHA-2 CHAIN (EHA-ALPHA-2) (ANTEN-NA PIGMENT PROTEIN, ALPHA-2 CHAIN) (FRAGMENT).//1.0:15:60//ECTOTHIORHODOSPIRA HALOPHILA.// P80101

F-HEMBA1001109//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//6.7e-37:102:82//HOMO SAPIENS (HU-MAN).//P39189

F-HEMBA1001121//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.036:49:46//HOMO SAPIENS (HU-MAN).//P08547

40 F-HEMBA1001122

F-HFMBA1001123

F-HEMBA1001133/HYPOTHETICAL 9.4 KD PROTEIN (ORF2).//0.86:29:41/FELINE IMMUNODEFICIENCY VI-RUS (ISOLATE SAN DIEGO) (FIV), AND FELINE IMMUNODEFICIENCY VIRUS (ISOLATE PETALUMA) (FIV).//

45 F-HEMBA1001137//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2).//2.0e-22:103:52//HOMO SA-PIENS (HUMAN).//P51523

F-HEMBA1001140//COLLAGEN ALPHA 4(IV) CHAIN PRECURSOR.//0.032:94:36//HOMO SAPIENS (HUMAN).// P53420

F-HEMBA1001172

F-HEMBA1001174//ADP-RIBOSYLATION FACTOR-LIKE PROTEIN 5.//2.9e-78:179:79//RATTUS NORVEGICUS (RAT).//P51646

F-HEMBA1001197/MAJOR PRION PROTEIN PRECURSOR (PRP) (PRP27-30) (PRP33-35C) (FRAGMENT).// 0.051:96:32//CERCOCEBUS ATERRIMUS, AND MACACA SYLVANUS (BARBARY APE).//Q95145

F-HEMBA1001208

55 F-HEMBA1001213

F-HEMBA1001226//FROTEASOME COMPONENT C8 (EC 3.4.99.46) (MACROPAIN SUBUNIT C8) (MULTICAT-ALYTIC ENDOPEPITIDASE COMPLEX SUBUNIT C8.9/1/1.5e-08:24:97/IHOMO SAPIENS (HUMAN)./P25788 F-HEMBA1001235//FIBRONECTIN (FN) (FRAGMENT)://0.756-038//ORYCTOLAGUS CUNICULUS (RABBIT).//

Q28749

F-HEMBA1001247//SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185).//0.00052:16:81//VOLVOX CART-FRI //P21997

F-HEMBA1001257//2-ARYLPROPIONYL-COA EPIMERASE (EC 5.-.-.).//1.6e-68:178:77//RATTUS NORVEGI-CUS (RAT).//P70473

F-HEMBA1001265//MANNAN ENDO-1.4-BETA-MANNOSIDASE A PRECURSOR (EC 3.2.1.78) (BETA- MAN-NANASE A) (1,4-BETA-D-MANNAN MANNANOHYDROLASE A),//0.67:23:60//PIROMYCES SP.//P55296 F-HEMBA1001281//HYPOTHETICAL 8.9 KD PROTEIN YCF34 (ORF76).//0.83:48:35//PORPHYRA PURPU-REA //P51229

F-HEMBA1001286//COMPLEMENT DECAY-ACCELERATING FACTOR PRECURSOR. //1.3e-07:185:29//CAVIA 10 PORCELLUS (GUINEA PIG).//Q60401

F-HEMBA1001289//METABOTROPIC GLUTAMATE RECEPTOR 3 PRECURSOR://0.00018;159:30//RATTUS NORVEGICUS (RAT)//P31422

F-HEMBA1001294

F-HEMBA1001299//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/5.3e-07:27:77//HOMO SAPIENS (HUMAN).// 15 P30105

F-HEMBA1001302//45 KD CALCIUM-BINDING PROTEIN PRECURSOR (STROMAL CELL-DERIVED FACTOR 4) (SDF-4).//3.3e-61:150:76//MUS MUSCULUS (MOUSE).//Q61112 F-HEMBA1001303

20 F-HEMBA1001310//HYPOTHETICAL PROTEIN KIAA0161.//2.7e-10:170:27//HOMO SAPIENS (HUMAN).// P50876

F-HEMBA1001319

F-HEMBA1001323

F-HEMBA1001326//HYPOTHETICAL 55.1 KD PROTEIN IN FAB1-PES4 INTERGENIC REGION //1 1e-39:144: 38//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43601

F-HEMBA1001327

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F-HEMBA1001330

F-HEMBA1001351//VESICLE-ASSOCIATED MEMBRANE PROTEIN/SYNAPTOBREVIN BINDING PROTEIN (VAP-33),//1.9e-37:155:46//APLYSIA CALIFORNICA (CALIFORNIA SEA HARE),//Q16943

F-HEMBA1001361//RUBREDOXIN (RD).//0.95:44:29//ALCALIGENES EUTROPHUS.//P31912 F-HEMBA1001375//AEROLYSIN REGULATORY PROTEIN. //0.013:45:33//AEROMONAS SOBRIA. //P09165 F-HEMBA1001377//SPERM PROTAMINE P1.//1.0:22:40//PLANIGALE MACULATA SINUALIS (COMMON PLAN-

IGALE).//O18746 F-HEMBA1001383//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN).//0.60:37:29//HUMAN IM-MUNODEFICIENCY VIRUS TYPE 1 (Z2/CDC-Z34 ISOLATE) (HIV-1).//P12506

F-HEMBA1001387//GTP-BINDING PROTEIN TC10.//6.6e-43:83:92//HOMO SAPIENS (HUMAN).//P17081 F-HEMBA1001388//HYPOTHETICAL PROTEIN KIAA0136 (FRAGMENT).//0.00088:46:45//HOMO SAPIENS (HUMAN).//Q14149

F-HEMBA1001391

F-HEMBA1001398//CLOACIN (EC 3.1.-.-) (RIBONUCLEASE).//1.0:59:37//ESCHERICHIA COLL.//P00645 40 F-HEMBA1001405//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.25:41:34//HOMO SAPIENS

F-HEMBA1001407//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONES CP3, CP4 AND CP5) ICON-TAINS: BASIC PEPTIDE IB-6: PEPTIDE P-HI.//4.0e-09:129:40//HOMO SAPIENS (HUMAN).//P04280

45 F-HEMBA1001411//HYPOTHETICAL 34.9 KD PROTEIN IN CYSJ-ENO INTERGENIC REGION (O313).//0.95:88: 31//ESCHERICHIA COLL//P55140

F-HEMBA1001413//SOX-12 PROTEIN (FRAGMENT),//0.95:46:32//MUS MUSCULUS (MOUSE),//Q04890 F-HEMBA1001415//HISTONE H5.//0.43:95:29//GALLUS GALLUS (CHICKEN).//P02259

F-HEMBA1001432//LANTIBIOTIC NISIN A PRECURSOR.//0.77:46:32//LACTOCOCCUS LACTIS (SUBSP. LAC-TIS) (STREPTOCOCCUS LACTIS).//P13068

F-HEMBA1001433//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.8e-09:132:31//NYCTICEBUS COU-CANG (SLOW LORIS), //P08548

F-HEMBA1001435//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.2e-31:84:77//HOMO SAPIENS (HUMAN),// P39189

55 F-HEMBA1001442

> F-HEMBA1001446//ANTIFREEZE PEPTIDE 4 PRECURSOR://0.71;41;39//PSEUDOPLEURONECTA AMERI-CANUS (WINTER FLOUNDER),//P02734

F-HEMBA1001450//PROLINE-RICH PROTEIN LAS17.//0.13:127:27//SACCHAROMYCES CEREVISIAE (BAK-

ER'S YEAST) //Q12446

F-HEMBA1001454//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.57:38:47//HANSENULA WINGEI (YEAST).//P48882

F-HEMBA1001455//CHEMOTAXIS PROTEIN CHEA (EC 2.7.3.-).//0.98:124:25//BORRELIA BURGDORFERI (LYME DISEASE SPIROCHETE).//Q44737

F-HEMBA1001463//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.8e-32:62:67//HOMO SAPIENS (HU-MAN) //P39194

F-HEMBA1001476//NUCLEOPORIN NUP159 (NUCLEAR PORE PROTEIN NUP159).//6.8e-09:252:36//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40477

F-HEMBA1001478

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F-HEMBA1001497//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.2e-33:105:72//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1001510//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//8.3e-37:54:81//HOMO SAPIENS (HUMAN).// P39189

15 F-HEMBA1001515//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//2.0e-63:223:57//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1001517

F-HEMBA1001522//TROPOMYOSIN ALPHA CHAIN, SMOOTH MUSCLE.//0.78:150:22//COTURNIX COTURNIX JAPONICA (JAPANESE QUAIL).//P49437

F-HEMBA1001526//PERIPLASMIC [FE] HYDROGENASE 1 (EC 1.18.99.1).//1.6e-06:130:29//CLOSTRIDIUM PASTEURIANUM.//P29166

F-HEMBA1001533//PROBABLE E5A PROTEIN.//0.73:35:37//HUMAN PAPILLOMAVIRUS TYPE 6A.//Q84296
F-HEMBA1001557//HYPOTHETICAL 17.1 KD PROTEIN IN PURS 3"REGION.//1.5e-07:99:36//SACCHAROMY-

CES CEREVISIAE (BAKER'S YEAST).//P38898 F-HEMBA1001566//HYPOTHETICAL PROTEIN BB0692.//0.91:27:44//BORRELIA BURGDORFERI (LYME DIS-

EASE SPIROCHETE).//051635 F-HEMBA1001569//SYNAPTOBREVIN 2 (VESICLE ASSOCIATED MEMBRANE PROTEIN 2) (VAMP-2).//2.2e-

50:110:95//HOMO SAPIENS (HUMAN), AND BOS TAURUS (BOVINE).//P19065 F-HEMBA1001570//IIII ALU SUBFAMILY SX WARNING ENTRY IIIII//3.3e-33:107:72//HOMO SAPIENS (HUMAN).//P39195

F-HEMBA1001579//RING CANAL PROTEIN (KELCH PROTEIN).//1.2e-14:111:39//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652

F-HEMBA1001581

F-HEMBA1001585

F-HEMBA1001589//PROBABLE DNA-BINDING PROTEIN (AGNOPROTEIN).//0.98:51:33//HUMAN ADENOVI-RUS TYPE 2.//P03263

F-HEMBA1001595//SEPTIN 2 HOMOLOG (FRAGMENT).//3.0e-124:274:85//HOMO SAPIENS (HUMAN).// Q14141

F-HEMBA1001608//RENAL SODIUM/DICARBOXYLATE COTRANSPORTER (NA(+)/DICARBOXYLATE COTRANSPORTER) //0.99:28:39//ORYCTOLAGUS CUNICULUS (RABBIT) //0.28615

F-HEMBA1001620//MYO-INOSITOL-1-PHOSPHATE SYNTHASE (EC 5.5.1.4) (IPS).//4.3e-45:222:46// SPIRODELA POLYRRHIZA.//P42803

F-HEMBA1001635//FIBRILLARIN.//0.10:72:38//CAENORHABDITIS ELEGANS.//Q22053

F-HEMBA1001636//PAIRED BOX PROTEIN PAX-8, ISOFORMS 8C/8D.//0.75:38:47//HOMO SAPIENS (HU-45 MAN).//Q09155

F-HEMBA1001640//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/14.7e-06:80:41//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1001647//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII-135) (TAFII-130) (TAFII-130).//0.075:165:32//HOMO SAPIENS (HUMAN).//000268

F-HEMBA1001651//GOLGIN-95.//6.8e-05:141:24//HOMO-SAPIENS (HUMAN).//Q08379

F-HEMBA1001655//PROLINE-RICH PROTEIN LAS17.//0.19:97:30//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//012446

F-HEMBA1001658/I/TETRAHYDROMETHANOPTERIN S-METHYLTRANSFERASE 12 KD SUBUNIT (EC 21.18 B) (NS-METHYLTETRAHYDROMETHANOPTERIN-COEX/YME M METHYLTRANSFERASE 12/K OSUB-UNIT)/I 0.29:44/METHANOBACTERIUM THERMOAUTOTROPHICUM (STRAIN MARBURG / DSM 2133)// 060773

F-HEMBA1001661//CELLULOSE COMPLEMENTING PROTEIN.//0.35:87:33//ACETOBACTER XYLINUM (ACETOBACTER PASTEURIANUS).//P37697

- F-HEMBA1001672//CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS).//2.7e-10:216:35//PLASMODIUM CYNOMOLGI (STRAIN BEROK).//P08672
- F-HEMBA1001675/I/NODULIN 20 PRECURSOR (N-20),//0.98:36:44//GLYCINE MAX (SOYBEAN),//P08960 F-HEMBA1001678//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//8.2e-13:62:64/HOMO SAPIENS (HUMAN),// P39195
- F-HEMBA1001681//HYPOTHETICAL 41.5 KD PROTEIN IN P6.5-VP48 INTERGENIC REGION (P40) (ORF3) (ORF102).//1.0:51:39//ORGYIA PSEUDOTSUGATA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//
- F-HEMBA1001702/IATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6).//0.017:54:37//TRYPANOSOMA BRU-CEI BRUCEI //P24499
- F-HEMBA1001709/HYPOTHETICAL 21.2 KD PROTEIN IN TOR2-MNN4 INTERGENIC REGION.//0.59:109:35// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36042
- F-HEMBA1001711
  F-HEMBA1001712//HYPOTHETICAL 6.9 KD PROTEIN IN 100 KD PROTEIN REGION.//0.54:44:34//HUMAN AD-
- 15 ENOVIRUS TYPE 41.//P23690
  F-HEMBA1001714//ATPASE\_INHIBITOR\_MITOCHONDRIAL\_PRECURSOR.//1.2e-19:60:75//RATTUS\_NOR-
  - VEGICUS (RAT).//Q03344

    F-HEMBA1001718//HYPOTHETICAL PROTEIN UL63.//1.0:54:37//HUMAN CYTOMEGALOVIRUS (STRAIN
- AD169).//P16820
  F-HEMBA1001723//HYPOTHETICAL 34.0 KD TRP-ASP REPEATS CONTAINING PROTEIN IN SIS1-MRPL2
- INTERGENIC REGION.//5.1-26:90:53//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P41318 F-HEMBA1001731/HYPOTHETICAL 16.6 KD PROTEIN.//0.71:49:32//AVIAN INFECTIOUS BURSAL DISEASE VIRUS (STRAIN 52/70) (IBDV).//P25221 F-HEMBA1001734
- 25 F-HEMBA1001744//SCY1 PROTEIN.//2.1e-11:182:29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P53009
  - F-HEMBA1001745//HYPOTHETICAL 11.6 KD PROTEIN IN NUT1-ARO2 INTERGENIC REGION PRECUR-SOR.//1.0:36:38//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53116
  - F-HEMBA1001746//PROTEIN-EXPORT MEMBRANE PROTEIN SECG HOMOLOG.//0.94:48:35//MYCOBACTE-RILIM LEPRAE //P38388
- F-HEMBA1001761

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- F-HEMBA1001781//ZINC FINGER PROTEIN 19 (ZINC FINGER PROTEIN KOX12) (FRAGMENT).//0.028:47:40//
  HOMO SAPIENS (HUMAN).//P17023
- F-HEMBA1001784/HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN CHROMOSOME X.//0.00068:32:46// CAENORHABDITIS ELEGANS .//Q11116
  - F-HEMBA1001791/METALLOTHIONEIN (MT)//1.034:35/PLEURONECTES PLATESSA (PLAICE)//P07216 F-HEMBA1001800//ZINC FINGER PROTEIN MFG-1 (ZINC FINGER PROTEIN 58) (FRAGMENT).//1.5e-14:60: 48/MUS MUSCULUS (MOUSE)//P16372
- F-HEMBA1001803
- 40 F-HEMBA1001804/IGLYCINE-RICH CELL WALL STRUCTURAL PROTEIN 1 PRECURSOR.//9.3e-17:56:57// ORYZA SATIVA (RICE).//P25074
  - F-HEMBA1001808//PARANEOPLASTIC ENCEPHALOMYELITIS ANTIGEN HUD HOMOLOG (HU-ANTIGEN D).//0.75:97:31//RATTUS NORVEGICUS (RAT).//009032
- F-HEMBA1001809//IMMEDIATE-EARLY PROTEIN IE180.//4.5e-11:206:36//PSEUDORABIES VIRUS (STRAIN INDIANA-FUNKHAUSER / BECKER) (PRV).//P11675
  - F-HEMBA1001815//60S RIBOSOMAL PROTEIN L37-B (YL27) (FRAGMENT).//0.34:30:30//SCHIZOSACCHA-ROMYCES POMBE (FISSION YEAST).//P06733
    - F-HEMBA1001819//ZINC FINGER PROTEIN 135.//2.6e-102:262:66//HOMO SAPIENS (HUMAN).//P52742 F-HEMBA1001820
- 50 F-HEMBA1001822//EPIDERMAL GROWTH FACTOR RECEPTOR SUBSTRATE SUBSTRATE 15 (PROTEIN EPS15).//1.2e-18:251:33//MUS MUSCULUS (MOUSE).//P42567
  - F-HEMBA1001824//KERATIN, ULTRA HIGH-SULFUR MATRIX PROTEIN (UHS KERATIN).//4.7e-11:124:37// OVIS ARIES (SHEEP).//P26372
- F-HEMBA1001835
- 55 F-HEMBA1001844//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//4.3e-14:36:63//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBA1001847//ZINC FINGER PROTEIN 29 (ZFP-29).//2.7e-36:135:51//MUS MUSCULUS (MOUSE).// Q07230

- F-HEMBA1001861
- F-HEMBA1001864//HEAT-STABLE ENTEROTOXIN A3/A4 PRECURSOR (STA3/STA4) (ST-IB) (ST-II).//1.0:31: 38/JESCHERICHIA COLI.//P07965
- F-HEMBA1001866//UDP-GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (EC 2.4.1.-) (DUGT).//9.7e-42:234:41//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q09332
- F-HEMBA1001869//HYPOTHETICAL 94.9 KD PROTEIN C22E12.11C IN CHROMOSOME I.//5.3e-13:65.47// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10362
  - F-HEMBA1001888//HYPOTHETICAL 11.4 KD PROTEIN (ORF1).//0.85:62:37//STREPTOMYCES FRADIAE.//
- 10 F-HEMBA1001896//DIMETHYLGLYCINE DEHYDROGENASE PRECURSOR (EC 1.5.99.2) (ME2GLYDH).//9.8e-20;250:29//RATTUS NORVEGICUS (RAT).//Q63342
  - F-HEMBA1001910//EUKARYOTIC TRANSLATION INITIATION FACTOR 4E (EIF-4E) (EIF4E) (MRNA CAP-BINDING PROTEIN) (EIF-4F 25 KD SUBUNIT).//0.94:44:38//CAENORHABDITIS ELEGANS.//O61955
- F-HEMBA1001912///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//8.7e-07:53:62//HOMO SAPIENS (HUMAN).//
  P30188
- F-HEMBA1001913//GCN20 PROTEIN.//1.8e-21:68:60//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  - F-HEMBA1001915//KLEE PROTEIN (KCRB3 PROTEIN).//0.94:64:21//ESCHERICHIA COLI.//Q52280
  - E-HEMBA1001018
- 20 F-HEMBA1001921
  - F-HEMBA1001939//CHLOROPLAST 50S RIBOSOMAL PROTEIN L24.//1.0:47:31//ODONTELLA SINENSIS.//
  - F-HEMBA1001940//!!! ALU SUBFAMILY J WARNING ENTRY !!!!/0.0017:31:77//HOMO SAPIENS (HUMAN).// P39188
- F-HEMBA1001942/IHIBERNATION-ASSOCIATED PLASMA PROTEIN HP-27 PRECURSOR (HIBERNATOR-SPECIFIC BLOOD COMPLEX, 27 KD SUBUNIT).//10.6777-28/I/TAMIAS ASIATICUS (CHIPMUNK).//Q06577 F-HEMBA1001945/I/HYPOTHETICAL 4.6 KD PROTEIN IN GP47-AGT INTERGENIC REGION (ORF E).//1.0:35:
  - 37//BACTERIOPHAGE T4.//P32269
    F-HEMBA1001950//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.43:18:44//DROSOPHILA YAKUBA
    (FRUIT FLY).//P03933
- 90 (FRUIT FLY)//P03933 F-HEMBA101960/HOMEOBOX PROTEIN HOX-C5 (HOX-3D) (CP11).//0.17:12:66//HOMO SAPIENS (HU-MAN) //O00444
  - F-HEMBA1001962/HYPOTHETICAL 9.0 KD PROTEIN IN ADH4 5'REGION.//1.0:30:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53056
  - F-HEMBA1001964

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- F-HEMBA1001967//HYPOTHETICAL PROTEIN UL61.//0.027:111:36//HUMAN CYTOMEGALOVIRUS (STRAIN AD169).//P16818
- F-HEMBA1001979

  F-HEMBA1001987//HYPOTHETICAL 11.2 KD PROTEIN (ORF117).//1.0.83:32//ORGYIA PSEUDOTSUGATA

  MULTICAPSID POLYHEDROSIS VIRUS (OPMMPV).//O10386
- F-HEMBA1001991//NEUROTOXIN 1 (TOXIN ATX-I).//0.99:31:45//ANEMONIA SULCATA (SNAKE-LOCKS SEA
  - F-HEMBA1002003//GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE, TESTIS-SPECIFIC (EC 1.2.1.12) (GAPDH).//5.5e-07:109:32//MUS MUSCULUS (MOUSE).//Q64467
- 45 F-HEMBA1002008
  - F-HEMBA1002018/IEC PROTEIN HOMOLOG 2 (FRAGMENT).//0.83:66:33//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//Q42377
    - F-HEMBA1002022//INSULIN.//1.0:59:32//SQUALUS ACANTHIAS (SPINY DOGFISH).//P12704
  - F-HEMBA1002035//MONOCYTIC LEUKEMIA ZINC FINGER PROTEIN.//8.3e-15:64:40//HOMO SAPIENS (HU-MAN).//Q92794
  - F-HEMBA1002039//HYPOTHETICAL PROLINE-RICH PROTEIN KIAA0269.//0.0070:70:40//HOMO SAPIENS (HUMAN).//Q92558
    - F-HEMBA1002049//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.5e-07:37:75//HOMO SAPIENS (HUMAN).//
- 55 F-HEMBA1002084
  - F-HEMBA1002092//SPT23 PROTEIN.//0.12:208:26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P35210
  - F-HEMBA1002100

- F-HEMBA1002102//ANKYRIN.//1.4e-12:106:35//MUS MUSCULUS (MOUSE).//Q02357
- F-HEMBA1002113//EARLY NODULIN 20 PRECURSOR (N-20).//0.073:155:32//MEDICAGO TRUNCATULA (BARREL MEDIC).//P93329
- F-HEMBA1002119//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.85:22:36//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643
- F-HEMBA1002125//GAG POLYPROTEIN [CONTAINS: CORE PROTEINS P15, P12, P30].//0.35:111:33//FELINE SARCOMA VIRUS (STRAIN SNYDER-THEILEN).//P03338
  - F-HEMBA1002139/HYPOTHETICAL 12.4 KD PROTEIN IN SEC17-QCR1 INTERGENIC REGION.//0.88:72:25// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38192
- F-HEMBA1002144

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- F-HEMBA1002150//THROMBOMODULIN (FETOMODULIN) (TM) (FRAGMENT).//4.8e-10:65:46//BOS TAURUS (BOVINE).//P06579
- F-HEMBA1002151/IA-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.24:146:28//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323
- F-HEMBA1002153//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3).//0.93:58:25//APIS MEL-LIFERA (HONEYBEE).//P34859
  - F-HEMBA1002160///!// ALU SUBFAMILY SP WARNING ENTRY !!!!//5.1e-21:94:65//HOMO SAPIENS (HUMAN).// P39193
  - F-HEMBA1002161//MYOSIN HEAVY CHAIN, CARDIAC MUSCLE BETA ISOFORM.//1.4e-51:180:56//SUS SCROFA (PIG).//P79293
  - F-HEMBA1002162//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//4.1e-40:102:75//HOMO SAPIENS (HU-MAN).//P39193
  - F-HEMBA1002166///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.8e-13:133:45//HOMO SAPIENS (HUMAN).// P39188
- 25 F-HEMBA1002177//ZINC FINGER PROTEIN 142 (KIAA0236) (HA4654).//0.0014:153:26//HOMO SAPIENS (HU-MAN).//P52746
  - F-HEMBA1002185
    F-HEMBA1002189/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/0.86:46:45//HOMO SAPIENS (HUMAN).//
- P39194

  F-HEMBA1002191//MALE SPECIFIC SPERM PROTEIN MST84DC.//0.037:14:57//DROSOPHILA MELA-NOGASTER (FRUIT FLY) //001644
  - F-HEMBA1002199
    - E-HEMBA1002204
    - F-HEMBA1002212//DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE
- 35 KINASE DSOR1 (EC 2.7.1.-) (DOWNSTREAM OF RAF) (MAPKK).//3.2e-13:201:30//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q24324
  - F-HEMBA1002215//TESTIN 2 (TES2) [CONTAINS: TESTIN 1 (TES1)].//1.1e-62:147:84//MUS MUSCULUS (MOUSE).//P47226
- F-HEMBA1002226//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.6e-26:168:44//HOMO SAPIENS (HUMAN).//
  40 P39188
  - F-HEMBA1002229//!!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!!/6.8e-18:68:72//HOMO SAPIENS (HU-MAN) //P39190
  - F-HEMBA1002237//EAMZP30-47 PROTEIN (FRAGMENT).//0.96:21:61//EIMERIA ACERVULINA.//P21959 F-HEMBA1002241//METALLOTHIONEIN (MT).//0.95:25:48//PARACENTROTUS LIVIDUS (COMMON SEA UR-
- 45 CHIN).//P80367 F-HEMBA1002253//METALLOTHIONEIN-II (MT-II).//0.97·27·48//MESOCRICETUS AURATUS (GOLDEN HAM-
  - F-HEMBA1002253/METALLOTHIONEIN-II (MT-II),//0.97:27:48//MESOCRICETOS AURATOS (GOLDEN HA STER),//P17808
    - F-HEMBA1002257
  - F-HEMBA1002265//MALE SPECIFIC SPERM PROTEIN MST84DC.//0.95:24:50//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01644
  - F-HEMBA1002267//NEURONAL PROTEIN 3.1 (P311 PROTEIN).//0.94:33:33//GALLUS GALLUS (CHICKEN).// Q90667
    - F-HEMBA1002270
  - F-HEMBA1002321//HYPOTHETICAL IMMUNITY REGION PROTEIN 14.//0.99:22:40//BACTERIOPHAGE PHI-
- 55 105.//P10437 F-HEMBA1002328

- F-HEMBA1002328
- F-HEMBA1002341//P53-BINDING PROTEIN 53BP2 (FRAGMENT).//3.7e-55:109:96//MUS MUSCULUS

- (MOUSE).//Q62415
- F-HEMBA1002348//PROBABLE E5 PROTEIN.//0.43:30:50//HUMAN PAPILLOMAVIRUS TYPE 35.//P27226
- F-HEMBA 1002349
- F-HEMBA1002363//CHROMOSOME ASSEMBLY PROTEIN XCAP-E.//5.7e-105:278:71//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P50533
- F-HEMBA1002381//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.3e-24:69:73//HOMO SAPIENS (HUMAN).// P39188
- F-HEMBA1002389//EARLY NODULIN 20 PRECURSOR (N-20).//0.16:110:31//MEDICAGO TRUNCATULA (BAR-REL MEDIC).//P93329
- F-HEMBA1002417//TIGHT JUNCTION PROTEIN ZO-1 (TIGHT JUNCTION PROTEIN 1).//2.6e-51:187:56//MUS MUSCULUS (MOUSE).//P39447
  - F-HEMBA1002419//PROLINE-RICH PEPTIDE P-B.//1.0:18:61//HOMO SAPIENS (HUMAN).//P02814
  - F-HEMBA1002430//HYPOTHETICAL 12.3 KD PROTEIN IN GAP1-NAP1 INTERGENIC REGION.//0.042:41:46// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36140
- 15 F-HEMBA1002439//CHLOROPLAST 50S RIBOSOMAL PROTEIN L27 (FRAGMENT).//0.99:47:29//CALYPTRO-SPHAERA SPHAEROIDEA.//P41548
  - F-HEMBA1002458//OVARIAN GRANULOSA CELL 13.0 KD PROTEIN HGR74.//4.1e-24:109:55//HOMO SAPIENS (HUMAN).//Q00994
- F-HEMBA1002460
- F-HEMBA1002462//SALIVARY PROLINE-RICH PROTEIN II-1 (FRAGMENT).//0.00025:80:30//HOMO SAPIENS (HUMAN).//P81489
  - F-HEMBA1002469//PUTATIVE TUMOR SUPPRESSOR LUCA15.//0.0012:110:33//HOMO SAPIENS (HUMAN).// P52756
- F-HEMBA1002475//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.37:106:33//MUS MUSCULUS (MOUSE).//P05143
- F-HEMBA1002477/IIII ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.3e-34:96:71//HOMO SAPIENS (HU-MAN) ///733194 F-HEMBA1002486
- F-HEMBA1002495/LIGHT-MEDIATED DEVELOPMENT PROTEIN DET1.//2.9e-31:110:39//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P48732
- F-HEMBA1002498//SFT2 PROTEIN.//1.0:54:35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  P38166
  - F-HEMBA1002503//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.0e-06:49:63//HOMO SAPIENS (HUMAN).//
- 35 F-HEMBA1002508/!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.6e-22:169:44//HOMO SAPIENS (HU-MAN).//P39195
  - F-HEMBA1002513//HYPOTHETICAL 89.8 KD PROTEIN F41H10.6 IN CHROMOSOME IV.//0.00017:79:35// CAENORHABDITIS ELEGANS.//Q20296
  - F-HEMBA1002515
- 40 F-HEMBA1002538//ATP SYNTHASE E CHAIN, MITOCHONDRIAL (EC 3.6.1.34).//1.0:53:37//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P81449
  - F-HEMBA1002542///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.7e-32:96:75//HOMO SAPIENS (HUMAN).// P39188
- F-HEMBA1002547//AGRIN PRECURSOR.//2.5e-100:218:80//RATTUS NORVEGICUS (RAT).//P25304
  45 F-HEMBA1002552//HEP27 PROTEIN (PROTEIN D).//9.5e-12:29:82//HOMO SAPIENS (HUMAN).//Q13268
  - F-HEMBA1002552//HEP27 PROTEIN (PROTEIN D).//9.5e-12:29:82//HOMO SAPIENS (HUMAN).//Q13268 F-HEMBA1002555//COLLAGEN ALPHA 1(III) CHAIN.//2.4e-15:207:36//BOS TAURUS (BOVINE).//P04258
  - F-HEMBA1002558//!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.0:34:50//HOMO SAPIENS (HUMAN).// P39193
  - F-HEMBA1002561/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.5e-05:49:46//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548
  - F-HEMBA1002569//SINGLE-STRANDED DNA-BINDING PROTEIN P12.//0.97:60:33//BACTERIOPHAGE
    - F-HEMBA1002583

- F-HEMBA1002590//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.6e-15:54:55//HOMO SAPIENS (HUMAN).//
- F-HEMBA1002592//HISTIDINE-RICH PROTEIN.//0.99:39:28//PLASMODIUM FALCIPARUM (ISOLATE FCM17 / SENEGAL).//P14586
  - F-HEMBA1002609//SSM4 PROTEIN.//1.9e-12:135:29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//

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F-HEMBA1002621//CYTOCHROME B6-F COMPLEX 3.5 KD SUBUNIT (CYTOCHROME B6-F COMPLEX SUB-UNIT 6).//1.0:20:55//ZEA MAYS (MAIZE).//P19445

F-HEMBA1002624//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).//
0.0035:124:33//NICOTIANA TABACUM (COMMON TOBACCO).//P13983

F-HEMBA1002628

F-HEMBA1002629//IMMEDIATE-EARLY PROTEIN IE180.//0.84:80:36//PSEUDORABIES VIRUS (STRAIN KAP-LAN) (PRV).//P33479

F-HEMBA1002645//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.8e-16:57:68//HOMO SAPIENS (HUMAN).//

F-HFMBA1002651

F-HEMBA1002659//CUTICLE COLLAGEN 2.//0.0077:77:38//CAENORHABDITIS ELEGANS.//P17656

F-HEMBA1002661//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG://2.3e-89:116:72//HOMO SAPIENS (HU-MAN)://P08547

15 F-HEMBA1002666//BETA CRYSTALLIN A4.//0.18:58:44//GALLUS GALLUS (CHICKEN).//P49152

F-HEMBA1002678

F-HEMBA1002679//GLUTAMIC ACID-RICH PROTEIN PRECURSOR.//5.7e-06:219:27//PLASMODIUM FALCI-PARUM (ISOLATE FC27 / PAPUA NEW GUINEA).//P13816

F-HEMBA1002688//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT).//1.1e-07:198:32//NEPHILA CLA-VIPES (ORB SPIDER).//P46804

F-HEMBA1002696//COLLAGEN ALPHA 1(VII) CHAIN PRECURSOR (LONG-CHAIN COLLAGEN) (LC COLLA-GEN).//0.16:158:33//HOMO SAPIENS (HUMAN).//Q02388

F-HEMBA1002703//HYPOTHETICAL BHLF1 PROTEIN.//0.78:147:29//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181

25 F-HEMBA1002712//11.2 KD PROTEIN (ORF 103).//0.029:75:34//BACTERIOPHAGE PF1.//P25133

E-HEMBA1002716//50S RIBOSOMAL PROTEIN I 28 //1 0:44:27//BACILLUS SUBTILIS //P37807

F-HEMBA1002728///!// ALU SUBFAMILY SX WARNING ENTRY !/!!//5.4e-18:56:75//HOMO SAPIENS (HUMAN).// P39195

F-HEMBA1002730//HYPOTHETICAL PROTEIN MJ0316.//0.097:84:35//METHANOCOCCUS JANNASCHII.//

30 Q57764 F-HEMBA1002742//APOLIPOPROTEIN C-III PRECURSOR (APO-CIII),//0.97:26:50//SUS SCROFA (PIG),//

P27917
F-HEMBA1002746//CALPHOTIN.//0.35:65:35//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q02910

F-HEMBA1002748//PLATELET GLYCOPROTEIN IB BETA CHAIN PRECURSOR (GP-IB BETA) (GPIBB).//1.0: 74:32//MUS MUSCULUS (MOUSE).//P56400

F-HEMBA1002750//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.0e-15:49:75//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1002768//HYPOTHETICAL 72.2 KD PROTEIN C12C2.05C IN CHROMOSOME II.//0.00036:197:26// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09746

F-HEMBA1002770//UTEROGLOBIN PRECURSOR (BLASTOKININ).//023:88:27//ORYCTOLAGUS CUNICU-LUS (RABBIT).//P02779

F-HEMBA1002777//HOMEOBOX PROTEIN HOX-A4 (HOX-1.4) (MH-3).//0.00018:67:43//MUS MUSCULUS (MOUSE).//P06798

F-HEMBA1002779//HYPOTHETICAL 17.6 KD PROTEIN IN NPR1-RPS3 INTERGENIC REGION.//0.70:30:53//
SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53880

F-HEMBA1002780//OLFACTORY RECEPTOR 3 (K10) (FRAGMENT).//1.0:31:45//MUS MUSCULUS (MOUSE).// Q60879

F-HEMBA1002794//HMG-Y RELATED PROTEIN B (SB16B PROTEIN) (FRAGMENT).//0.0044:66:37//GLYCINE MAX (SOYBEAN).//010370

F-HEMBA1002801

F-HEMBA1002810//HYPOTHETICAL 25.9 KD PROTEIN AH6.3 IN CHROMOSOME II.//0.0033:116:31// CAENORHABDITIS ELEGANS.//Q09202

F-HEMBA1002816//HYPOTHETICAL 47.1 KD PROTEIN C9G1.13C IN CHROMOSOME I.//1.0e-17:68:48// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O14308

55 F-HEMBA1002818//FIBULIN-2 PRECURSOR.//2.1e-27:92:44//MUS MUSCULUS (MOUSE).//P37889

F-HEMBA1002826//DNA-BINDING PROTEIN 65 (PROTEIN GP65).//0.28:46:34//BACTERIOPHAGE T4.// P16012

F-HEMBA1002833

- F-HEMBA1002850//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:27:37//METRIDIUM SENILE (BROWN SEA ANEMONE).//O47493
- F-HEMBA1002863//PHOTOSYSTEM | REACTION CENTRE SUBUNIT IV (PHOTOSYSTEM | 8.1 KD PROTEIN) (P30 PROTEIN) (PSI-E).//0.84:37:43//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//P12975
- F-HEMBA1002876//OCTAPEPTIDE-REPEAT PROTEIN T2.//0.74:58:34//MUS MUSCULUS (MOUSE).//Q06666 F-HEMBA1002886
  - F-HEMBA1002896//HOMEOBOX PROTEIN HOX-B3 (HOX-2G) (HOX-2.7).//4.7e-05:84:35//HOMO SAPIENS (HUMAN).//P14651
  - F-HEMBA1002921//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN).//0.21:42:42//HUMAN IMU-NODEFICIENCY VIRUS TYPE 1 (RF/HAT ISOLATE) (HIV-1).//P05908
  - F-HEMBA1002924//EC PROTEIN HOMOLOG 2 (FRAGMENT),//0.85:75:22//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS),//Q42377
  - F-HEMBA1002934//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.4e-31:92:72//HOMO SAPIENS (HUMAN).// P39188
- 15 F-HEMBA1002935//GASTRULA ZINC FINGER PROTEIN XLCGF58.1 (FRAGMENT).//7.7e-06:187:29//XENO-PUS LAEVIS (AFRICAN CLAWED FROG).//P18730
  - F-HEMBA1002937//SUPPRESSOR PROTEIN SRP40.//0.00031:150:24//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32583
  - F-HEMBA1002939//ANKYRIN R (ANKYRINS 2.1 AND 2.2) (ERYTHROCYTE ANKYRIN).//5.2e-25:225:33//HO-MO SAPIENS (HUMAN).//P16157
  - F-HEMBA1002944

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- F-HEMBA1002951//TRICHOHYALIN.//0.0011:220:24//HOMO SAPIENS (HUMAN).//007283
- F-HEMBA1002954//PROBABLE E8 PROTEIN.//0.98:49:32//BOVINE PAPILLOMAVIRUS TYPE 4.//P08352
- F-HEMBA1002968//ACCESSORY GLAND PEPTIDE PRECURSOR (PARAGONIAL PEPTIDE B).//0.93:41:34//
  DROSOPHILA SECHELLIA (FRUIT FLY).//O18417
  - F-HEMBA1002970//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.00010:35:62//HOMO SAPIENS (HU-MAN).//P39193
- F-HEMBA1002971/I/INSULIN./1.0:31:35/I/HYDROLAGUS COLLIEI (SPOTTED RATFISH) (RACIFIC RATFISH), AND CHIMAERA MONSTROSA (RABBIT FISH).//P09536 F-HEMBA1002973/I/CAMP-DEPENDENT 3,5-CY-CLIC PHOSPHODIESTERASE 48 (EC 3.14.17) (DPDE4)//3.0e-2963:100//RATTUS NORVEGICIE (RAT)//
- P14646
  F-HEMBA1002997//HYPOTHETICAL 106.5 KD PROTEIN IN CTT1-PRP31 INTERGENIC REGION.//1.0e-08: 211:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53253
  - F-HEMBA1002999/SUPPRESSOR PROTEIN SRP40.//0.026:175:23//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583
    - ER'S YEAST)//P32583 F-HEMBA1003021/!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!/1.3e-36:102:70//HOMO SAPIENS (HU-MAN)//P39194
    - F-HEMBA1003033//HYPOTHETICAL 23.1 KD PROTEIN CY277.20C.//0.029:75:29//MYCOBACTERIUM TU-BERCIJI OSIS //P71779
- 40 F-HEMBA1003034//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//6.3e-23:144:46//HOMO SAPIENS (HU-MAN) //P39192
  - F-HEMBA1003035//HYPOTHETICAL 13.3 KD PROTEIN IN AROD-COMER INTERGENIC REGION.//0.99:55: 30//BACILLUS SUBTILIS.//P54457
- F-HEMBA1003037//DNA-BINDING PROTEIN INHIBITOR ID-4.//0.17:42:40//HOMO SAPIENS (HUMAN).//
  45 P47928
  - F-HEMBA1003041//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.69:28:46//HO-MO SAPIENS (HUMAN).//P30808
    - F-HEMBA1003046//MITOCHONDRIAL PROCESSING PROTEASE BETA SUBUNIT PRECURSOR (EC 3.4.24.64) (BETA-MPP) (P-52).//7.9e-124:253:96//HOMO SAPIENS (HUMAN).//O75439
    - F-HEMBA1003064//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3).//0.25:263:22//TRYPANO-SOMA BRUCEI BRUCEI.//P04540
      - F-HEMBA1003067/H/YPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//4.1e-05: 189:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214 F-HEMBA1003071//CUTICLE COLLAGEN 40 //6.0e-07:126:38//CAENORHABDITIS ELEGANS.//P34804
- 55 F-HEMBA1003077//FIBROMODULIN PRECURSOR (FM) (COLLAGEN-BINDING 59 KD PROTEIN)//2.4e-12: 139:34//HOMO SAPIENS (HUMAN).//Q06828
  - F-HEMBA1003078//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE].//7.2e-05;60:40//MUS MUSCULUS (MOUSE).//P11369

- F-HEMBA1003079//PROTEIN Q300.//0.0012:16:87//MUS MUSCULUS (MQUSE).//Q02722
- F-HEMBA1003083//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//3.3e-32:95:75//HOMO SAPIENS (HUMAN).//
- F-HEMBA1003086
- F-HEMBA1003096//PROTAMINE IA (IRIDINE IA).//0.36:20:40//SALMO IRIDEUS (RAINBOW TROUT).//P02328 F-HEMBA1003098/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/4.4e-09:43:72//HOMO SAPIENS (HUMAN).// P30488
  - F-HEMBA1003117//PUTATIVE CUTICLE COLLAGEN C09G5.5.//1.0:88:38//CAENORHABDITIS ELEGANS.//
- 10 F-HEMBA1003129//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3).//0.61:63:25//APIS MEL-LIFERA (HONEYBEE).//P34859
  - F-HEMBA1003133//COLLAGEN ALPHA 2(VIII) CHAIN (ENDOTHELIAL COLLAGEN) (FRAGMENT).//0.48:79: 37//HOMO SAPIENS (HUMAN).//P25067
- F-HEMBA1003136/MANNOSE-1-PHOSPHATE GUANYLTRANSFERASE (EC 2.7.7.13) (ATF-MANNOSE-1PHOSPHATE GUANYLYLTRANSFERASE) (NDP-HEXOSE PYROPHOSPHORYLASE).//3.6e-25:190:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P41940
  - F-HEMBA1003142
    F-HEMBA1003148/HYPOTHETICAL 56.4 KD PROTEIN IN RPL30-CWH41 INTERGENIC REGION PRECUR-SOR //0.088-171-23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P53189
- 20 F-HEMBA1003166//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.8e-13:54:66//HOMO SAPIENS (HUMAN).//
- F-HEMBA1003175//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.015:147:
  - 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P53214
    F-HEMBA1003179//PROBABLE TRNA (5-METHYLAMINOMETHYL-2-THIOURIDYLATE)-METHYLTRANS-FERASE (EC. 2.1.1.81)//2.6e-51:164-47//BACILLUS SUBTILIS //035020
- F-HEMBA1003197
  F-HEMBA1003199/HOMEOBOX PROTEIN HOX-44 (HOX-1D) (HOX-1.4\//0.00049:83:38/HOMO SAPIENS
  - (HUMAN).//Q00056 F-HEMBA1003202//SPERM PROTAMINE P1.//0.98:53:28//PLANIGALE GILESI (FLAT-SKULLED MARSUPIAL
- MOUSE).//018747
  F-HEMBA1003204///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!//5 2e-22:42:80//HOMO SAPIENS (HU-
  - MAN).//P39194
    F-HEMBA1003212//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.6e-18:74:71//HOMO SAPIENS (HUMAN).//
  - P39193
    F-HEMBA1003220/!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//3.3e-18:56:78//HOMO SAPIENS (HU-MAN)//P39194
    - F-HEMBA1003222//HYPOTHETICAL 37.5 KD PROTEIN IN GNTR-HTPG INTERGENIC REGION.//0.0018:159: 27//BACILLUS SUBTILIS //P46327
- F-HEMBA1003229//DIHYDRODIPICOLINATE SYNTHASE 1 PRECURSOR (EC 4.2.1.52) (DHDPS).//1.0:85:28//
  TRITICUM AESTIVUM (WHEAT).//P24846
- F-HEMBA1003235//TROPOMYOSIN://8.3e-07:109:33//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q02088
- F-HEMBA1003250

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- F-HEMBA1003257//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//1.5e-07:27:74//OWENIA FUSI-
- 45 FORMIS.//P21260
  - F-HEMBA1003273 F-HEMBA1003276
    - F-HEMBA1003278
  - F-HEMBA1003281//HOMEOBOX PROTEIN HOX-A4 (CHOX-1.4).//0.0053:116:36//GALLUS GALLUS (CHICK-EN).//P17277
- F-HEMBA1003266//DNA-DIRECTED RNA POLYMERASE SUBUNIT N (EC 2.7.7.6).//0.96:37:35//SULFOLOBUS ACIDOCAL DARIUS.//P39472
  - F-HEMBA1003291//5'-AMP-ACTIVATED PROTEIN KINASE, CATALYTIC ALPHA-2 CHAIN (EC 2.7.1.-) (AMPK ALPHA-2 CHAIN) (FRAGMENT).//3.3e-15:68:39//SUS SCROFA (PIG).//Q28948
- 59 F-HEIMBA1003298//PULMONARY SURFACTANT-ASSOCIATED PROTEIN B (SP-B) (6 KD PROTEIN) (PULMO-NARY SURFACTANT-ASSOCIATED PROTEIDLIPD (SP-12) (9-12) (

- F-HEMBA1003309//HYPOTHETICAL 7.9 KD PROTEIN.//0.69:54:37//VACCINIA VIRUS (STRAIN WR), AND VACCINIA VIRUS (STRAIN COPENHAGEN).//P04306
- F-HEMBA1003314//MIXED LINEAGE KINASE 2 (EC 2.7.1.-) (FRAGMENT).//2.3e-06:143:22//HOMO SAPIENS (HUMAN).//002779
- F-HEMBA1003322///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.5e-30:53:77//HOMO SAPIENS (HU-MAN).//P39194
  - F-HEMBA1003327
  - F-HEMBA1003328/TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN) (FRAGMENT).//0.53:21:42// HUMAN IMMUNODEFICIENCY VIRUS TYPE 1 (BH5 ISOLATE) (HIV-1).//P04612
- F-HEMBA1003330/LONG NEUROTOXIN 3 (TOXIN VN2).//1.0:26:34//DENDROASPIS POLYLEPIS POLYLEPIS (BLACK MAMBA).//P25667
  - F-HEMBA1003348/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/6.5e-09:56:66//HOMO SAPIENS (HU-MAN).//P39194
  - F-HEMBA1003369//ANTER-SPECIFIC PROLINE-RICH PROTEIN APG PRECURSOR.//0.0042:97:36//ARABI-DOPSIS THALIANA (MOUSE-EAR CRESS).//P40602
  - DOPSIS THALIANA (MOUSE-EAR CRESS).//P40602
    F-HEMBA1003370//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.0e-18:99:53//HOMO SAPIENS (HUMAN).//
    - F-HEMBA1003373

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- F-HEMBA1003376//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//4.7e-16:60:75//HOMO SAPIENS (HUMAN).//
- F-HEMBA1003380//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.8e-10:50:68//HOMO SAPIENS (HUMAN).//
- F-HEMBA1003384
- F-HEMBA1003395//PROBABLE E5 PROTEIN.//0.62:64:29//HUMAN PAPILLOMAVIRUS TYPE 16.//P06927
- 25 F-HEMBA1003402//HYPOTHETICAL 12.0 KD PROTEIN IN TUB1-CPR3 INTERGENIC REGION PRECURSOR.// 0.89:74:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04521
  - F-HEMBA1003403//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT).//0.0010: 69:33//RATTUS NORVEGICUS (RAT).//P10164
- F-HEMBA1003408//WEB1 PROTEIN (PROTEIN TRANSPORT PROTEIN SEC31).//4.8e-06:93:25//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P38968
- F-HEMBA1003417//PROCOLLAGEN ALPHA 1(II) CHAIN PRECURSOR [CONTAINS: CHONDROCALCIN].//
  0.0021:140:34//MUS MUSCULUS (MOUSE).//P28481
  - F-HEMBA1003418//SPLICING FACTOR, ARGININE/SERINE-RICH 4 (PRE-MRNA SPLICING FACTOR SRP75).//1.7e-14:188:33//HOMO SAPIENS (HUMAN).//Q08170
- F-HEMBA1003433//DNA REPAIR PROTEIN XRS2.//1.0:88:35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P33301
  - F-HEMBA1003447//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT).//0.0061: 69:33//RATTUS NORVEGICUS (RAT).//P10164
- F-HEMBA1003461//SPIDROIN 1 (DRAGLINE SILK FIBROIN 1) (FRAGMENT).//2.3e-09:239:33//NEPHILA CLA-40 VIPES (ORB SPIDER).//P19837
  - F-HEMBA1003463//METALLOTHIONEIN-A (MTA) (FRAGMENT).//1.0:40:35//SPHAERECHINUS GRANULARIS (PURPLE SEA URCHIN).//Q26497
  - F-HEMBA1003480//FUSARIC ACID RESISTANCE PROTEIN FUSB.//0.0043:96:32//BURKHOLDERIA CEPACIA (PSEUDOMONAS CEPACIA) //P24127
- 45 F-HEMBA1003528//36.4 KD PROLINE-RICH PROTEIN://6.4e-15:167:33//LYCOPERSICON ESCULENTUM (TOMATO)://Q00451
  - F-HEMBA1003531//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.2e-18:56:78//HOMO SAPIENS (HUMAN).// P39189
  - F-HEMBA1003538//COMPLEMENT C1R COMPONENT PRECURSOR (EC 3.4.21.41).//2.5e-28:136:47//HOMO SAPIENS (HUMAN).//P00736
    - F-HEMBA1003545//INSULIN GENE ENHANCER PROTEIN ISL-2 (ISLET-2).//9.2e-105:217:85//RATTUS NOR-VEGICUS (RAT).//P50480
    - F-HEMBA1003548
- F-HEMBA1003555//HYPOTHETICAL 31.9 KD PROTEIN IN BET1-PAN1 INTERGENIC REGION.//8.7e-57:180: 55//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40558
  - F-HEMBA1003556//HYPOTHETICAL 19.2 KD PROTEIN IN COX-REP INTERGENIC REGION (ORF5) (ORF21).// 0.53:97:25//BACTERIOPHAGE HP1.//P51706
  - F-HEMBA1003560//GUANINE NUCLEOTIDE-BINDING PROTEIN G(I)/G(S)/G(O) GAMMA-2 SUBUNIT (G GAM-

MA-I).//1.8e-32:71:100//BOS TAURUS (BOVINE).//P16874

F-HEMBA1003568//ZINC-FINGER PROTEIN RFP (RET FINGER PROTEIN).//4.1e-19:126:31//HOMO SAPIENS (HUMAN).//P14373

F-HEMBA1003569//METASTASIS-ASSOCIATED PROTEIN MTA1.//3.9e-83:143:74//HOMO SAPIENS (HU-MAN).//Q13330

F-HEMBA1003571//HYPOTHETICAL 8.7 KD PROTEIN (READING FRAME D).//1.0:64:25//STAPHYLOCOCCUS

F-HEMBA1003579//CYTOTOXIN 1 (CYTOTOXIN V-II-1) (TOXIN V(II)1).//1.0:41:29//NAJA MELANOLEUCA (FOREST COBRA) (BLACK-LIPPED COBRA).//P01448

10 F-HEMBA1003581//TALIN.//3.7e-36:52:98//MUS MUSCULUS (MOUSE).//P26039

F-HEMBA1003591//CHLOROPLAST 28 KD RIBONUCLEOPROTEIN PRECURSOR (28RNP).//1.6e-05:91:31//
NICOTIANA SYLVESTRIS (WOOD TOBACCO).//P19682

F-HEMBA1003595/HYPOTHETICAL 12.0 KD PROTEIN IN DST1-HEM2 INTERGENIC REGION.//1.0:55:32// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53182

15 F-HEMBA1003597

F-HEMBA1003598/T-CELL RECEPTOR BETA CHAIN PRECURSOR (ANA 11)//4.9e-10:85:41//ORYCTOLAGUS CUNICULUS (RAPRIT) //P06333

F-HEMBA1003615//PUTATIVE MINOR COAT PROTEIN (ORF43).//0.086:10:70//BACTERIOPHAGE PHI-LF.//

F-HEMBA1003617//HYPOTHETICAL 36.8 KD PROTEIN C26A3.16 IN CHROMOSOME I.//4.4e-13:58:48// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10169

F-HEMBA1003621//LONG NEUROTOXIN 1 (NEUROTOXIN A).//0.096:40:37//OPHIOPHAGUS HANNAH (KING COBRA) (NAJA HANNAH).//P01387

F-HEMBA1003622

F-HEMBA1003630

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F-HEMBA1003637//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.4e-13:47:74//HOMO SAPIENS (HUMAN).//

F-HEMBA1003640//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.87:25:64//HOMO SAPIENS (HUMAN).//

30 F-HEMBA1003645//HYPOTHETICAL 40.4 KD TRP-ASP REPEATS CONTAINING PROTEIN C14B1.4 IN CHRO-MOSOME III.//1.8e-10:157:26//CAENORHABDITIS ELEGANS.//Q17963

F-HEMBA1003646//SERINE-ARGININE PROTEIN 55 (SRP55) (ENHANCER OF DEFORMED) (52-KD BRACK-ETING PROTEIN) (852 PROTEIN).//4.96-05:207:27//DROSOPHILA MELANOGASTER (FRUIT FLY).//P26686 F-HEMBA1003656

F-HEMBA1003662//PROLINE-RICH PEPTIDE P-B.//0.57:17:52//HOMO SAPIENS (HUMAN).//P02814

F-HEMBA1003667//!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!/6.0e-16:43:72//HOMO SAPIENS (HU-MAN)//P39194

F-HEMBA1003679

F-HEMBA1003680//PUTATIVE AMINOPEPTIDASE ZK353.6 IN CHROMOSOME III (EC 3.4.11.-).//3.9e-08:137: 27//CAENORHABDITIS ELEGANS //P34629

F-HEMBA1003684/IZINC FINGER PROTEIN 151 (POLYOMAVIRUS LATE INITIATOR PROMOTER BINDING PROTEIN) (LP-1) (ZINC FINGER PROTEIN Z13).//2.1e-20:127:40//MUS MUSCULUS (MOUSE).//Q60821

F-HEMBA1003690//HYPOTHETICAL PROTEIN KIAA0288 (HA6116).//3.0e-85:201:78//HOMO SAPIENS (HU-MAN).//P56524

45 F-HEMBA1003692//CELL DIVISION CONTROL PROTEIN 1.//0.13:69:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40986

F-HEMBA1003711//CARCINOEMBRYONIC ANTIGEN PRECURSOR (CEA) (MECONIUM ANTIGEN 100) (CD66E ANTIGEN).//0.021:153:26//HOMO SAPIENS (HUMAN).//P06731

F-HEMBA1003714//ABAECIN.//0.99:34:32//BOMBUS PASCUORUM.//P81463

50 F-HEMBA1003715

F-HEMBA1003720//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//5.4e-34:155:56//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1003725//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.3e-27:181:41//HOMO SAPIENS (HU-MAN).//PD8547

F-HEMBA1003729//PTB-ASSOCIATED SPLICING FACTOR (PSF).//0.0037:103:33//HOMO SAPIENS (HU-MAN)//P323/6

F-HEMBA1003733//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//5.0e-54:210:58//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1003742//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.066;72:33//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643

F-HEMBA1003758

F-HEMBA1003760//HYPOXIA-INDUCIBLE FACTOR 1 ALPHA (HIF-1 ALPHA) (ARNT INTERACTING PRO-TEIN).//1.5e-51:220:52//MUS MUSCULUS (MOUSE).//Q61221

F-HEMBA1003773

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F-HEMIDA 1003773

F-HEMBA1003783 F-HEMBA1003784

F-HEMBA1003799//SHORT NEUROTOXIN 1 (TOXIN AA C).//0.95:27:37//ACANTHOPHIS ANTARCTICUS (COMMON DEATH ADDER).//P01434

F-HEMBA1003803//GAG PÓLYPROTEIN [CONTAINS: CORE PROTEINS P15, P12, P30].//0.46:96:34//FELINE SARCOMA VIRUS (STRAIN SNYDER-THEILEN).//P03338

F-HEMBA1003804//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.019:30:50//HO-MO SAPIENS (HUMAN).//P30808

15 F-HEMBA1003805//HYPOTHETICAL 75.0 KD PROTEIN B0280.11 IN CHROMOSOME III.//1.8e-20:109:47// CAENORHABDITIS ELEGANS //P42083

F-HEMBA1003807

F-HEMBA1003827//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//2.1e-09:23:78//OWENIA FUSI-FORMIS //P21260

F-HEMBA1003836//MOB1 PROTEIN (MPS1 BINDER 1).//2:0e-31:134:52//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40484

F-HEMBA1003838///!/I ALU SUBFAMILY SC WARNING ENTRY !!!!//3.9e-22:39:76//HOMO SAPIENS (HUMAN).// P39192

F-HEMBA1003856

25 F-HEMBA1003864/HYPOTHETICAL 39.4 KD PROTEIN IN MET1-SIS2 INTERGENIC REGION.//1.5e-15:194: 30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36151

F.-HEMBA1003866//PROTEIN A39 //0 0027.72:33//VACCINIA VIRUS (STRAIN COPENHAGEN).//P21062 F.-HEMBA1003879//80 KD NUCLEAR CAP BINDING PROTEIN (NCBP 80 KD SUBUNIT) (CBP80).//2.9e-16:22: 100//HOMO SAPIENS (HUMAN).//009161

30 F-HEMBA1003880//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.99:39:38//FELIS SILVESTRIS CATUS (CAT).//P48896

F-HEMBA1003885//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//3.5e-28:47:76//HOMO SAPIENS (HUMAN).// P39193

F-HEMBA1003893//HYPOTHETICAL 27.8 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//1.7e-57: 215:51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53215

F-HEMBA1003902

F-HEMBA1003908

F-HEMBA1003926//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.3e-10:60:63//HOMO SAPIENS (HUMAN).//

40 F-HEMBA1003937//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//8.1e-29:68:64//HOMO SAPIENS (HU-MAN) //P39194

F-HEMBA1003939//PROTEIN Q300.//0.0025;24:62//MUS MUSCULUS (MOUSE).//Q02722

F-HEMBA1003942//EXCITATORY INSECT TOXIN BJXTR-IT PRECURSOR (BJ-XTRIT).//0.084;67:31//BUTHO-TUS JUDAICUS (SCORPION) (HOTTENTOTTA JUDAICA).//P56637

45 F-HEMBA10039\$0//HYPOTHÉTICAL 8.1 KD PROTEIN IN SPEA-METK INTERGENIC REGION (O71).//0.95:26: 34//ESCHERICHIA COLI.//P46878

F-HEMBA1003953//ZINC FINGER PROTEIN MFG-1 (ZINC FINGER PROTEIN 58) (FRAGMENT).//2.5e-17:89: 46//MUS MUSCULUS (MOUSE).//P16372

F-HEMBA1003958/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.2e-23:43:76//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1003959

F-HEMBA1003976//HYPOTHETICAL PROTEIN KIAA0076 (HA0936).//0.99:88:28//HOMO SAPIENS (HUMAN).// Q14999

F-HEMBA1003978//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.98:19:57//HOMO SAPIENS (HUMAN).//P22531

F-HEMBA1003985//LYSYL-TRNA SYNTHETASE (EC 6.1.1.6) (LYSINE-TRNA LIGASE) (LYSRS) (FRAG-MENT).//1.0:40:32//MYCOBACTERIUM LEPRAE.//P46861

F-HEMBA1003987//HYPOTHETICAL PROTEIN UL66.//0.27:65:33//HUMAN CYTOMEGALOVIRUS (STRAIN

AD169) //P16822

F-HEMBA1003989/MALE SPECIFIC SPERM PROTEIN MST84DB.//5.2e-05:64:40//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643

F-HEMBA1004000//PROTEIN Q300.//0.00042:17:82//MUS MUSCULUS (MQUSE).//Q02722

F-HEMBA1004011//ALPHA-TYPE CALCITONIN GENE-RELATED PEPTIDE PRECURSOR (CGRP-1).//0.47: 106:32//HOMO SAPIENS (HUMAN).//P06881

F-HEMBA1004012//ATP SYNTHASE PROTEIN 9, MITOCHONDRIAL (EC 3.6.1.34) (LIPID-BINDING PROTEIN).//0.96:36:33//PARAMECIUM TETRAURELIA.//P16001

F-HEMBA1004015//HYPOTHETICAL 29.3 KD PROTEIN B0280.6 IN CHROMOSOME III.//0.00018:90:34// CAENORHABDITIS ELEGANS.//P41997

F-HEMBA1004024//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//5.1e-34:75:80//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1004038

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F-HEMBA1004042

15 F-HEMBA1004045//40S RIBOSOMAL PROTEIN S27A.//1.0:20:55//ASPARAGUS OFFICINALIS (GARDEN AS-PARAGUS).//P31753

F-HEMBA1004048//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//1.3e-06:158:35//MUS MUSCULUS (MOUSE).//P05143

F-HEMBA1004049//32 KD HEAT SHOCK PROTEIN (4-1 PROTEIN).//0.098:106:32//DICTYOSTELIUM DISCOI-DEUM (SLIME MOLD).//P54658

F-HEMBA1004055//HYPOTHETICAL PROTEIN HI0258/259.//0.87:133:23//HAEMOPHILUS INFLUENZAE.// P43974

F-HEMBA1004056///!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//3.3e-25:39:64//HOMO SAPIENS (HU-MAN)//P39191

25 F-HEMBA1004074///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.9e-08:35:68//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1004086

F-HEMBA1004097//IMMEDIATE-EARLY PROTEIN IE4 (IE68) (FRAGMENT)//0.71:95:35//HERPES SIMPLEX VI-RUS (TYPE 2).//P14379

30 F-HEMBA1004111/IIII ALU SUBFAMILY J WARNING ENTRY !!!!/4.7e-26:84:64/HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1004131//SEPTIN 2 HOMOLOG (FRAGMENT).//2.8e-34:108:63//HOMO SAPIENS (HUMAN).// Q14141

F-HEMBA1004132/HYPOTHETICAL PROTEIN HI1738./I1.0:44:34/HAEMOPHILUS INFLUENZAE./IP44300
F-HEMBA1004133/HYPOTHETICAL 8.5 KD PROTEIN CY274.40C./I0.89:21:57//MYCOBACTERIUM TUBER-CULOSIS.//C10826

F-HEMBA1004138//EARLY NODULIN 75 (N-75) (NGM-75) (FRAGMENT).//0.016:39:41//MEDICAGO SATIVA (ALFALFA).//P11728

F-HEMBA1004143//CYTOCHROME C OXIDASE POLYPEPTIDE VIII PRECURSOR (EC 1.9.3.1).//0.93:34:29// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P04039

F-HEMBA1004146//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.63:52:36//HOMO SAPIENS (HUMAN).// P02811

F-HEMBA1004150//METALLOTHIONEIN-II (MT-II).//1.0:20:45//MUS MUSCULUS (MOUSE).//P02798

F-HEMBA1004164//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!!//3.0e-13:57:71//HOMO SAPIENS (HUMAN).// P39195

F-HEMBA1004168/V-TYPE SODIUM ATP SYNTHASE SUBUNIT F (EC 3,6.1.34) (NA(+)-TRANSLOCATING AT-PASE SUBUNIT F).//0.00035-90:34/ENTEROCOCCUS HIRAE.//P43437
F-HEMBA1004199//HYPOTHETICAL HELICASE K12H4.8 IN CHROMOSOME III.//5.1e-14:115:31//

F-HEMBA1004199//HYPOTHETICAL HELICASE K12H4.8 IN CHROMOSOME III.//5.1e-14:115:31/.
CAENORHABDITIS ELEGANS.//P34529

F-HEMBA1004200

F-HEMBA1004202/YPT1-RELATED PROTEIN 1.//2.5e-24:96:52//SCHIZOSACCHAROMYCES POMBE (FIS-SION YEAST).//P11620

F-HEMBA1004203///!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.2e-09:48:64//HOMO SAPIENS (HUMAN).// P39193

F-HEMBA1004207//HYPOTHETICAL 8.7 KD PROTEIN IN RPL22-RPL23 INTERGENIC REGION (ORF70).// 0.98:51:33//ASTASIA LONGA (EUGLENOPHYCEAN ALGA).//P34779

F-HEMBA1004225//METALLOTHIONEIN-II.//1.0:30:33//CANDIDA GLABRATA (YEAST) (TORULOPSIS GLABRATA).//P15114

F-HEMBA1004227//PUTATIVE PROTEIN PHOSPHATASE 2C (EC 3.1.3.16) (PP2C) (KIAA0015).//5.9e-06:109: 33//HOMO SAPIENS (HUMAN).//P49593

F-HEMBA1004238I/VERY HYPOTHETICAL XYLU PROTEIN./I0.98:39:38/IESCHERICHIA COLI.//P05056 F-HEMBA1004241//SOX-13 PROTEIN (FRAGMENT)./I0.66:36:38//MUS MUSCULUS (MOUSE).//Q04891

F-HFMBA1004246

F-HEMBA1004248/I/INSULIN-INDUCED GROWTH RESPONSE PROTEIN CL-6 (IMMEDIATE-EARLY PROTEIN CL-6) //1.0e-43:98:84//RATTUS NORVEGICUS (RAT) //008755

F-HEMBA1004264//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT).//0.014:160:28//NEPHILA CLA-VIPES (ORB SPIDER).//P46804

F-HEMBA1004267//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.8e-52:56:83//HOMO SAPIENS (HUMAN).// P39189

F-HEMBA1004272

F-HEMBA1004274//HYPOTHETICAL 13.0 KD PROTEIN F59B2.10 IN CHROMOSOME III.//0.00084:33:54//
CAENORHABDITIS ELEGANS.//P34485

15 F-HEMBA1004275//HYPOTHETICAL 56.5 KD PROTEIN IN CAJ1-HOM3 INTERGENIC REGION)/9.3e-06:125: 27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40034

F-HEMBA1004276/IBETA-ADAPTIN 1 (PLASMA MEMBRANE ADAPTOR HAZ/AP2 ADAPTIN BETA SUBUNIT) (CLATHRIN ASSEMBLY PROTEIN COMPLEX 2 BETA LARGE CHAIN) (AP105A).//3.7e-30:239:32//HOMO SA-PIENS (HUMAN)./010567

F-HEMBA1004286//CUTICLE COLLAGEN 34.//0.0027:71:38//CAENORHABDITIS ELEGANS.//P34687 F-HEMBA1004289//PTR3 PROTEIN (SSY3 PROTEIN).//1.0:76:28//SACCHAROMYCES CEREVISIAE (BAK-

F-HEMBA1004299/PFR3 PROTEIN (SS13 PROTEIN)/II.0.76.20/SACCHAROWITCES CEREVISIAE (BAN-ER'S YEAST).//P43606 F-HEMBA1004299//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.075:58:39/HO-

MO SAPIENS (HUMAN).//P30808 F-HEMBA1004306//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//0.020:132:30//ORGYIA PSEUDOTSUGATA

MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV),//O10341
F-HEMBA1004312//EARLY PROTEIN 173R.//0.99:65:32//AFRICAN SWINE FEVER VIRUS (STRAIN BA71V)

(ASFV).//P27946
F-HEMBA1004321//ZINC FINGER PROTEIN 90 (ZFP-90) (ZINC FINGER PROTEIN NK10).//4.3e-43:133:44//

30 MUS MUSCULUS (MOUSE).//Q61967 F-HEMBA1004323

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F-HEMBA1004327//SMALL PROLINE-RICH PROTEIN 2-1.//0.027.48.43//HOMO SAPIENS (HUMAN).//P35326
F-HEMBA1004330//HOMEOBOX PROTEIN ENGRAILED-1 (HU-EN-1).//0.46.70:34//HOMO SAPIENS (HU-MAN).//COGATOR.

F-HEMBA1004334//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//7.7e-05:83:34//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1004335///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.0e-24:41:80//HOMO SAPIENS (HUMAN).// P39195

F-HEMBA1004341//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//2.8e-06:148:35//MUS MUSCULUS (MOUSE).//P05143

F-HEMBA1004353//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.2e-29:57:80//HOMO SAPIENS (HUMAN).// P39195

F-HEMBA1004354//CHL1 PROTEIN.//0.017:40:40//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//

45 F-HEMBA1004356

F-HEMBA1004366/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.00045:49:46//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1004372//VERY HYPOTHETICAL 20.6 KD PROTEIN C56F8.15 IN CHROMOSOME I.//1.0:125:28// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10263

F-HEMBA1004389//HYPOTHETICAL 113.1 KD PROTEIN IN PRE5-FET4 INTERGENIC REGION.//0.76:170:25// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04893

F-HEMBA1004394

F-HEMBA1004396//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.2e-10:72:51//HOMO SAPIENS (HU-MAN).//P08547

55 F-HEMBA1004405

F-HEMBA1004408//PEPTIDYL-PROLYL CIS-TRANS ISOMERASE 10 (EC 5.2.1.8) (PPIASE) (ROTAMASE) (CY-CLOPHILIN-10).//2.7e-29:146:48//CAENORHABDITIS ELEGANS.//P52017

F-HEMBA1004429/!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//0.0019:47:59//HOMO SAPIENS (HU-

MAN).//P39191

- F-HEMBA1004433//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.1e-20:47:68//HOMO SAPIENS (HUMAN).//
- F-HEMBA1004460//!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//6.2e-64:134:69//HOMO SAPIENS (HU-MAN).//P39193
  - F-HEMBA1004461//METALLOTHIONEIN-LIKE PROTEIN 1.//1.0:39:35//PISUM SATIVUM (GARDEN PEA).// P20830
- F-HEMBA1004479//HYPOXIA-INDUCIBLE FACTOR 1 ALPHA (HIF-1 ALPHA) (ARNT INTERACTING PRO-TEIN).//9.7e-43:101:48//MUS MUSCULUS (MOUSE).//Q61221
- 10 F-HEMBA1004482//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34).//1.0:41:36//CANDIDA GLABRATA (YEAST) (TORULOPSIS GLABRATA).//P05040
  - F-HEMBA1004499//TUBULIN BETA CHAIN.//0.00021:55:36//CAENORHABDITIS ELEGANS.//P52275
  - F-HEMBA1004502 F-HEMBA1004506//HYPOTHETICAL PROTEIN ORF-1137.//5.3-11:119:35//MUS MUSCULUS (MOUSE).//
- 15 P11260 F-HEMBA1004507//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//0.00072:90:37//HOMO
  - F-HEMBA 1004501/1/SFLICEUSUME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).10.00072:30:37/IHOMO SAPIENS (HUMAN)./Q15428 F-HEMBA 1004509/HYPOTHETICAL 52.2 KD PROTEIN IN MPR1-GCN20 INTERGENIC REGION./6.3e-28:169:
  - 42//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P43589
    F-HEMBA1004534/ENDOTHELIAL ACTIN-BINDING PROTEIN (ABP-280) (NONMUSCLE FILAMIN) (FILAMIN
- 1).//1.3e-80:226:66//HOMO SAPIENS (HUMAN).//P21333 F-HEMBA1004538//HYPOTHETICAL PROTEIN MJ0764.//0.96:28:35//METHANOCOCCUS JANNASCHII.//
  - F-HEMBA1004542//METALLOTHIONEIN (MT).//0.78:36:41//GADUS MORHUA (ATLANTIC COD).//P51902
- 25 F-HEMBA1004554

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- F-HEMBA1004560//HYPOTHETICAL PROTEIN KIAA0281 (HA6725).//4.2e-15:56:69//HOMO SAPIENS (HU-MAN)//002556
- F-HEMBA1004573//CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS).//0.65:31:58//PLASMODIUM BERGHEI.//P06915
- 30 F-HEMBA1004577//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//3.9e-08:35:80//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBA1004586//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!/6.6e-08:64:54//HOMO SAPIENS (HU-MAN)./P39194
  - F-HEMBA1004596//HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN C (HNRNP C) (HNRNP CORE PROTEIN C) (FRAGMENT).//0.00057:88:31//RATTUS NORVEGICUS (RAT).//P17132
  - F-HEMBA1004604//COLLAGEN ALPHA 2(XI) CHAIN PRECURSOR (FRAGMENT).//0.045:37:45//MUS MUSCU-LUS (MOUSE).//Q64739
  - F-HEMBA1004610//!!! ALU SUBFAMILY J WARNING ENTRY !!!!/4.3e-11:73:54//HOMO SAPIENS (HUMAN).// P39188
- 40 F-HEMBA1004617
- F-HFMBA1004629
  - F-HEMBA1004631//HYPOTHETICAL 7.8 KD PROTEIN IN WAPA-LICT INTERGENIC REGION.//1.0:36:38//BA-CILLUS SUBTILIS://P42303
- F-HEMBA1004632//PHOTOSYSTEM I REACTION CENTRE SUBUNIT X PRECURSOR (LIGHT-HARVESTING 8.0 KD POLYPEPTIDE).//0.86:48:35//SYNECHOCOCCUS ELONGATUS NAEGELI.//P20453
  - F-HEMBA1004637/HYPOTHETICAL 83.6 KD PROTEIN R05D3.2 IN CHROMOSOME III.//1.7e-32:159:42//
    CAENORHABDITIS ELEGANS.//P34535
    - F-HEMBA1004638//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//2.8e-06:50:46//OWENIA FUSI-FORMIS.//P21260
- 50 F-HEMBA1004666/TOXIN S6C4.//1.0:36:30//DENDROASPIS JAMESONI KAIMOSAE (EASTERN JAMESON'S MAMBA).//P25682
  - F-HEMBA1004669//SPLICING FACTOR, ARGININE/SERINE-RICH 4 (PRE-MRNA SPLICING FACTOR SRP75).//1.6e-12:105:42//HOMO SAPIENS (HUMAN).//Q08170
- F-HEMBA1004670//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//2.5e-06:62:45//HOMO SAPIENS (HU-55 MAN).//P02452
  - F-HEMBA1004672//HYPOTHETICAL PROTEIN MJ0437.//0.95:37:29//METHANOCOCCUS JANNASCHII.//
    - F-HEMBA1004693//MYOSIN HEAVY CHAIN, NONMUSCLE TYPE B (CELLULAR MYOSIN HEAVY CHAIN,

TYPE B) (NMMHC-B)://0.00035:217:23//HOMO SAPIENS (HUMAN)://P35580

F-HEMBA1004697/IMMUNOGLOBULIN G BINDING PROTEIN H PRECURSOR (PROTEIN H).//0.058:118:30// STREPTOCOCCUS PYOGENES.//P50470

F-HEMBA1004705///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//6.8e-09:43:72//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1004709/!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//8.8e-18:50:84//HOMO SAPIENS (HUMAN).// P39189

F-HEMBA1004711//ETS-RELATED PROTEIN 71 (ETS TRANSLOCATION VARIANT 2).//0.0027:148:30//HOMO SAPIENS (HUMAN).//000321

F-HEMBA1004725//CUTICLE COLLAGEN 2.//0.0051:41:41//CAENORHABDITIS ELEGANS.//P17656

F-HEMBA1004730//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.4e-22:210:37//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1004733//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.7e-07:50:62//HOMO SAPIENS (HUMAN).// P39188

15 F-HEMBA1004734/I/UBIQUITIN-CON, IUGATINO ENZYME E2-18 KD (EC 6.3.2.19) (UBIQUITIN- PROTEIN LIGASE) (UBIQUITIN CARRIER PROTEIN) (PM42).//9.99-39:143:52//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS)/P42743

F-HEMBA1004736/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.1e-60:210:61//HOMO SAPIENS (HU-MAN).//P08547

F-HFMBA1004748

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F-HEMBA1004751//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.8e-20:88:63//HOMO SAPIENS (HUMAN).// P39188

F-HEMBA1004752//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//0.0043:126:34// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437

25 F-HEMBA1004753//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/7.8e-28:47:78//HOMO SAPIENS (HUMAN).// P39193

F-HEMBA1004756/HYPOTHETICAL 53.3 KD PROTEIN IN HXT8-CAN1 INTERGENIC REGION.//0.22:77:27// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39981 F-HFMBA1004758

F-HEMIDA 1004/36

90 F-HEMBA1004763/HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT),//1.1e-06:58:43//OWENIA FUSI-FORMIS //P21260 F-HEMBA1004768/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.7e-65:298:53/HOMO SAPIENS (HU-

F-HEMBA1004768//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.7e-65:298:53//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBA1004770

F-HEMBA1004771

F-HEMBA1004776//GRANULIN 1.//0.78:28:42//CYPRINUS CARPIO (COMMON CARP).//P81013

F-HEMBA1004778

F-HEMBA1004795//CDC4-LIKE PROTEIN (FRAGMENT).//6.9e-20:74:63//HOMO SAPIENS (HUMAN).//P50851 F-HEMBA1004803/kINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.4e-22:58:86//HOMO SAPIENS (HUMAN).//P508547

F-HEMBA1004806//HYPOTHETICAL 24.3 KD PROTEIN IN PSBH-RPL11 INTERGENIC REGION (ORF182).//
0.72:75:33//CYANOPHORA PARADOXA.//P48324

F-HEMBA1004807

F-HEMBA1004816

45 F-HEMBA1004820//HEMOLYMPH TRYPSIN INHIBITOR A (BPI-TYPE) (FRAGMENT).//1.0:50:38//MANDUCA SEXTA (TOBACCO HAWKMOTH) (TOBACCO HORNWORM).//P26226

F-HEMBA1004847//SIGNAL RECOGNITION PARTICLE 68 KD PROTEIN (SRP68).//3.0e-76:171:91//CANIS FA-MILIARIS (DOG).//Q00004

F-HEMBA1004850//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//3.0e-05:64:43//BOS TAURUS (BO-VINE).//P25508

F-HEMBA1004863/TOXIN C13S1C1 PRECURSOR.//0.38:52:30//DENDROASPIS ANGUSTICEPS (EASTERN GREEN MAMBA).//P18329

F-HEMBA1004864//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN) (FRAGMENT).//0.89:24:50//
HUMAN IMMUNODEFICIENCY VIRUS TYPE 1 (BH5 ISOLATE) (HIV-1).//P04612

55 F-HEMBA1004865

F-HEMBA1004880

F-HEMBA1004889//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//0.66:23:47//HOMO SAPIENS (HUMAN).//P22532

- F-HEMBA1004900
- F-HEMBA1004900
- F-HEMBA1004918/ICHLOROPLAST 30S RIBOSOMAL PROTEIN S8 (FRAGMENT), //0.56:37:32//SPINACIA OL-ERACEA (SPINACH), //P09597
- 5 F-HEMBA1004923//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3,5e-24:44:68//HOMO SAPIENS (HUMAN).//
  - F-HEMBA1004929//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.97:39:38//STRONGYLOCENTROTUS
  - PURPURATUS (PURPLE SEA URCHIN).//P15997
    F-HEMBA1004930//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.6e-15:64:59//HOMO SAPIENS (HU-
  - MAN).//P08547
    F-HEMBA1004933//VASODILATOR-STIMULATED PHOSPHOPROTEIN (VASP).//0.34:58:41//HOMO SAPIENS
    - (HUMAN).//P50552 F-HEMBA1004934
    - F-HEMBA1004944
- 15 F-HEMBA1004954/NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 3 (EC 1.6.5.3).//0.58:78:30//PARA-MECIUM TETRAURELIA.//P15579
  - F-HEMBA1004956//HYPOTHETICAL 18.8 KD PROTEIN (ORF4).//0.98:57:31//PARAMECIUM TETRAURELIA.// P15605
  - F-HEMBA1004960//HYPOTHETICAL 12.6 KD PROTEIN-(ORFJ) (RETRON EC67).//1.0:58:27//ESCHERICHIA COLI //P21324
  - F-HEMBA1004972
    - F-HEMBA1004973//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.90:55:30//HOMO SAPIENS (HUMAN).//P22531
    - F-HEMBA1004977
- 25 F-HEMBA1004978

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- F-HEMBA1004980//MOTILIN PRECURSOR.//0.088:79:31//MACACA MULATTA (RHESUS MACAQUE).//018811 F-HEMBA1004983//10 KD CHAPERONIN (PROTEIN CPN10) (PROTEIN GROES).//0.87:51:31//BUCHNERA APHIDICIA 4//058176
- F-HEMBA1004995//MYOCYTE-SPECIFIC ENHANCER FACTOR 2B (SERUM RESPONSE FACTOR-LIKE PRO-TEIN 2) (XMEF2) (RSRFR2).//0.17:52:40//HOMO SAPIENS (HUMAN).//Q02080
- F-HEMBA1005008//METALLOTHIONEIN (MT).//1.0:52:32//CRASSOSTREA VIRGINICA (EASTERN OYS-TER) //P23038
  - F-HEMBA1005009//ACTIN.//3.5e-27:171:38//CANDIDA ALBICANS (YEAST).//P14235
  - F-HEMBA1005019//HYPOTHETICAL PROTEIN HI1222.//0.13:58:31//HAEMOPHILUS INFLUENZAE.//P44129 F-HEMBA1005029//P2Y PURINOCEPTOR 5 (P2Y5) (PURINERGIC RECEPTOR 5) (6H1).//0.76:72:31//GALLUS
    - GALLUS (CHICKEN).//P32250
      F-HEMBA1005035//HOMEOBOX PROTEIN HB9.//0.0086:60:40//HOMO SAPIENS (HUMAN).//P50219
    - F-HEMBA1005039//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//0.47:49:32//HOMO SAPIENS (HUMAN).//P22532
- 40 F-HEMBA1005047//RAS-RELATED PROTEIN RAB-24 (RAB-16).//1.5e-19:39:100//MUS MUSCULU
  - F-HEMBA1005050//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.074:34:44//BOS TAURUS (BOVINE).// P25508
  - F-HEMBA1005062
- 45 F-HEMBA1005066//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//2.1e-44:126:65//HOMO SAPIENS (HU-MAN).//P08547
  - F-HEMBA1005075//SUPPRESSOR PROTEIN SRP40.//0.35:96:31//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583
  - F-HEMBA1005079/!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//3.6e-20:75:64//HOMO SAPIENS (HU-MAN) //P39191
- F-HEMBA1005083//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.00015:72:34//BOS TAURUS (BO-VINE).//P25508
  - F-HEMBA1005101//HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN 27C (HNRNP 48) (HRP48.1).//
    4.8e-10:176:25//DROSOPHILA MELANOGASTER (FRUIT FLY).//P48809
- 55 F-HEMBA1005113
  - F-HEMBA1005123//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.6e-24:99:60//HOMO SAPIENS (HU-MAN).//P39194
  - F-HEMBA1005133//HYPOTHETICAL 13.5 KD PROTEIN IN MOB1-SGA1 INTERGENIC REGION.//0.11:22:54//

- SACCHAROMYCES CEREVISIAE (BAKER'S YEAST). //P40490
- F-HEMBA1005149//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.7e-16:59:71//HOMO SAPIENS (HUMAN).// P39188
- F-HEMBA1005152/IGENOME POLYPROTEIN 2 (CONTAINS: HELPER COMPONENT PROTEINASE (EC 3.4.22.-) (HC-PRO); 70 KD PROTEIN].//1.0:77:27//BARLEY YELLOW MOSAIC VIRUS (JAPANESE STRAIN II-1) (BAYMV).//I/O01207
  - F-HEMBA1005159//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3).//0.40:53:33//APIS MEL-LIFERA (HONEYBEE).//P34859
- F-HEMBA1005185//MYOSIN IB HEAVY CHAIN.//0.011:58:48//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P34092
  - F-HEMBA1005201//HYPOTHETICAL 56.6 KD PROTEIN C16C9.03 IN CHROMOSOME I.//3.9e-67:241:53// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09817
- F-HEMBA1005202//SIGNAL RECOGNITION PARTICLE 68 KD PROTEIN (SRP68).//3.8e-124:257:95//CANIS FAMILIARIS (DOG).//Q00004
- 15 F-HEMBA1005206//CUTICLE COLLAGEN 1.//0.010:118:33//CAENORHABDITIS ELEGANS.//P08124
  - F-HEMBA1005219//PTB-ASSOCIATED SPLICING FACTOR (PSF).//0.99:85:40//HOMO SAPIENS (HUMAN).// P23246
    - F-HEMBA1005223//HYPOTHETICAL GENE 1.05 PROTEIN.//0.31:75:28//BACTERIOPHAGE T3.//P07715
    - F-HEMBA1005232//HYPOTHETICAL 7.8 KD PROTEIN.//0.99:48:29//VACCINIA VIRUS (STRAIN WR), AND VACCINIA VIRUS (STRAIN COPENHAGEN).//P20544
    - F-HEMBA1005241//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.4e-28:138:55//HOMO SAPIENS (HU-MAN)./P39193
      - F-HEMBA1005244//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.014:39:41//HOMO SAPIENS (HUMAN).//P22531
- 25 F-HEMBA1005251//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.55:15:46//DICENTRARCHUS LABRAX (EUROPEAN SEA BASS).//Q36362
  - F-HEMBA1005252//EC PROTEIN HOMOLOG (ZINC-METALLOTHIONEIN CLASS II).//0.088:33:42//ZEA MAYS (MAIZE).//F43401
    F-HEMBA1005724
- 30 F-HEMBA1005275/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/0.96:42:45//HOMO SAPIENS (HUMAN).// P39188
  - F-HEMBA1005293//PROBABLE COATOMER BETA' SUBUNIT (BETA'-COAT PROTEIN) (BETA'-COP).//0.55:98: 30//CAENORHABDITIS ELEGANS.//Q20168
  - F-HEMBA1005296/MUCIN 2 PRECURSOR (INTESTINAL MUCIN 2).//0.095:75:34//HOMO SAPIENS (HU-MAN).//Q02817
    - F-HEMBA1005304//!!!! ALU SUBFAMILY SB WARNING ENTRY!!!!/5.4e-33:103:74//HOMO SAPIENS (HU-MAN).//P39189
    - F-HEMBA1005311//PERIOD CLOCK PROTEIN (FRAGMENT).//0.99:45:31//DROSOPHILA SALTANS (FRUIT FLY).//Q04536
- 40 F-HÉMBA1005314//HYPOTHETICAL 6.3 KD PROTEIN T19C3.3 IN CHROMOSOME III.//0.98:30:30// CAENORHABDITIS ELEGANS.//Q10009
  - F-HEMBA1005315//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.1e-05:35:51//HOMO SAPIENS (HU-MAN).//P08547
- F-HEMBA1005318/IOLFACTORY RECEPTOR-LIKE PROTEIN COR8 (FRAGMENT).//IO.57:44:38//GALLUS
  45 GALLUS (CHICKEN).//Q98913
  F-HEMBA1005331/IMMEDIATE-EARLY PROTEIN IE180.//IO.57:106:33//PSEUDORABIES VIRUS (STRAIN IN-
  - DIANA-FUNKHAUSER / BECKER) (PRV).//P11675 F-HEMBA1005338//CARTIAGE MATRIX PROTEIN PRECURSOR (MATRILIN-1).//1.8e-55;199:59//GALLUS
  - F-HEMBA1005338//CARTIAGE MAIRIX PROTEIN PRECURSOR (MAIRILIN-1).//1.8e-55:199:59//GALLU
    GALLUS (CHICKEN).//P05099
  - F-HEMBA1005353//CHLOROPLAST 30S RIBOSOMAL PROTEIN S17.//0.88:33:36//PORPHYRA PURPUREA.// P51305
    - F-HEMBA1005359//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//1.1e-88:255:48//HOMO SA-PIENS (HUMAN).//P51522
- F-HEMBA1005367//ALPHA-AMYLASE INHIBITOR AAI.//1.0:25:40//AMARANTHUS HYPOCHONDRIACUS (PRINCE'S FEATHER).//P80403
  - F-HEMBA1005372

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F-HEMBA1005374///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.0e-34:92:75//HOMO SAPIENS (HU-MAN).//P39194

- F-HEMBA1005382//APOLIPOPROTEIN C-II (APO-CII).//0.99:39:33//BOS TAURUS (BOVINE).//P19034 F-HEMBA1005389//HYPOTHETICAL 70.0 KD PROTEIN IN DNAK 3'REGION (ORF4).//0.82:164:31//LACTO-
- COCCUS LACTIS (SUBSP. LACTIS) (STREPTOCOCCUS LACTIS).//P42377
- F-HEMBA1005394//HYPOTHETICAL 8.9 KD PROTEIN IN IEO-IE1 INTERGENIC REGION.//0.98:44:38// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41703
- F-HEMBA1005403//SPERM HISTONE P2 PRECURSOR (PROTAMINE MP2).//0.066:64:29//MUS MUSCULUS (MOUSE).//P07978
  - F-HEMBA1005408//50S RIBOSOMAL PROTEIN L33.//0.77:32:25//BACILLUS SUBTILIS.//Q06798
- F-HEMBA1005410//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE
- (EC 2.7.7.49): ENDONUCLEASEJ./0.006538:52/JMUS MUSCULUS (MOUSE).//P11369 F-HEMBA1005411//TOXIN S4C8.//0.16:46:28/JDENDROASPIS JAMESONI KAIMOSAE (EASTERN JAMES-ON'S MAMBA)//P25683
  - F-HEMBA1005423//CYCLIN-DEPENDENT KINASE 6 INHIBITOR (P18-INK6) (CYCLIN-DEPENDENT KINASE 4 INHIBITOR C) (P18-INK4C).//4.3e-09:29:96//HOMO SAPIENS (HUMAN).//P42773
- 15 F-HEMBA1005426/TOXIN C10S2C2.//0.99:49:34//DENDROASPIS ANGUSTICEPS (EASTERN GREEN MAM-BA).//P25684
  - F-HEMBA1005443///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.9e-16:78:60//HOMO SAPIENS (HUMAN).// P39188
  - F-HEMBA1005447//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.99:57:31//DASYPUS NOVEMCINCTUS (NINE-BANDED ARMADILLO).//021329
  - F-HEMBA1005468/INADH-UBIQUINONE OXIDOREDUCTASE CHAIN 1 (EC 1.6.5.3) (FRAGMENTS).//0.68:41: 31//ARTEMIA SALINA (BRINE SHRIMP).//P19040
    - F-HEMBA1005469
- F-HEMBA1005472//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.5e-39:142:70//HOMO SAPIENS (HU-MAN).//P08547
- F-HEMBA1005474//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//5.8e-10:44:68//HOMO SAPIENS (HU-
  - F-HEMBA1005475//U1 SMALL NUCLEAR RIBONUCLEOPROTEIN 70 KD (U1 SNRNP 70 KD) (SNRP70).//9.2e-14:179:33//HOMO SAPIENS (HUMAN).//P08621
- 30 F-HEMBA1005497

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- F-HEMBA1005500//60S RIBOSOMAL PROTEIN L37.//0.11:53:33//SCHISTOSOMA MANSONI (BLOOD FLUKE).//044125
- F-HEMBA1005506
- F-HEMBA1005508
- 35 F-HEMBA1005511//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.5e-30:92:73//HOMO SAPIENS (HUMAN).//
  - F-HEMBA1005513//MALES-ABSENT ON THE FIRST PROTEIN (EC 2.3.1.-).//2.0e-39:95:61//DROSOPHILA MELANOGASTER (FRUIT FLY).//002193
  - F-HEMBA1005517//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//2.1e-06:56:44//MUS MUSCULUS
- 40 (MOUSE),//P05142 F-HEMBA1005518//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS),//5.8e-05:192:33//BOS TAURUS (BO-
  - F-HEMBA1005520//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/2.0e-18:87:57//HOMO SAPIENS (HUMAN).//
- F-HEMBA1005526//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!/5.1e-22:77:54//HOMO SAPIENS (HU-
- MAN).//P39191 F-HEMBA1005528//CCR4-ASSOCIATED FACTOR 1 (CAF1).//1.2e-81:157:98//MUS MUSCULUS (MOUSE).//
  - F-HEMBA1005530//POLLEN ALLERGEN AMB P 5-A PRECURSOR (AMB P V-A).//0.98:19:47//AMBROSIA PSI-
- 50 LOSTACHYA (WESTERN RAGWEED).//P43174 F-HEMBA1005548//TRANSCRIPTION FACTOR MAF1.//1.4e-72:137:97//RATTUS NORVEGICUS (RAT).//
  - F-HEMBA1005552//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.7e-29:47:78//HOMO SAPIENS (HUMAN).//
- 55 F-HEMBA1005558/HYPOTHETICAL 25.6 KD PROTEIN IN ABF2-CHL12 INTERGENIC REGION.//1.6e-20:202: 30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04272
  - F-HEMBA1005568

P54842

F-HEMBA1005570//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 6 (EC 1.6.5.3).//1.0:80:31//

CAENORHABDITIS ELEGANS //P24885

- F-HEMBA1005576//TRANSMEMBRANE PROTEIN SEX PRECURSOR.//8.5e-58:152:75//HOMO SAPIENS (HU-MAN).//P51805
- F-HEMBA1005577//KERATIN, HIGH-SULFUR MATRIX PROTEIN, B2A.//0.98:57:36//OVIS ARIES (SHEEP).// P02438
- F-HEMBA1005581//SLIT PROTEIN PRECURSOR.//1.1e-62:254:41//DROSOPHILA MELANOGASTER (FRUIT FLY).//P24014
- F-HEMBA1005582//DYNACTIN, 150 KD ISOFORM (150 KD DYNEIN-ASSOCIATED POLYPEPTIDE) (DP-150) (DAP-150) (P150-GLUED).//0.0091:189:29//RATTUS NORVEGICUS (RAT).//P28023
- 10 F-HEMBA1005583//HYPOTHETICAL 41.2 KD PROTEIN IN CPS REGION (ORF7).//0.83:119:23//KLEBSIELLA PNEUMONIAE ///048453
  - F-HEMBA1005588//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.9e-17:108:53//HOMO SAPIENS (HUMAN).//
  - F-HEMBA1005593//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//0.23:24:54//HOMO SAPIENS (HUMAN).//P22532
  - F-HEMBA1005595//DYNEIN HEAVY CHAIN, CYTOSOUC (DYHC).//2.7e-39:257:39//DICTYOSTELIUM DISCOI-DEUM (SLIME MOLD).//P34036
    - F-HEMBA1005606

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- F-HEMBA1005609///!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//3.2e-20:27:96//HOMO SAPIENS (HUMAN).//
- F-HEMBA1005616//LATE CONTROL GENE B PROTEIN (GPB).//0.48:51:33//BACTERIOPHAGE 186.//P08711
  F-HEMBA1005617/IMITOTIC MAD2 PROTEIN.//1.2e-06:137:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//PAI088
- F-HEMBA1005627//HYPOTHETICAL 17.1 KD PROTEIN IN PUBS 3'REGION.//0.18:100:32//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P38898
- F-HEMBA1005631
  - F-HEMBA1005632//HYPOTHETICAL 7.4 KD PROTEIN.//0.32:59:32//VACCINIA VIRUS (STRAIN WR).//P04309 F-HEMBA1005634//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.6e-14:93:58//HOMO SAPIENS (HUMAN).// P30188
- 30 F-HEMBA1005666//HYPOTHETICAL PROTEIN KIAA0129.//2.1e-05:126:25//HOMO SAPIENS (HUMAN).// Q14142
  - F-HEMBA1005670
    - F-HEMBA1005679//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//2.2e-08:40:72//HOMO SAPIENS (HU-MAN).//P08547
- F-HEMBA1005680//SMALL PROLINE-RICH PROTEIN 2-1.//0.015:19:47//HOMO SAPIENS (HUMAN).//P35326 F-HEMBA1005685
  - F-HEMBA1005699/IEPHRIN-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) (LERK-8) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3)//4.2e-38/98/81/HOMO SAPIENS (HUMAN)//0.15768
- #0 F-HEMBA1005705//PROTEIN Q300.//0.11:23:56//MUS MUSCULUS (MOUSE).//Q02722
  - F-HEMBA1005717
  - F-HEMBA1005732//BACTENECIN 7 PRECURSOR (BAC7).//0.22:55:41//OVIS ARIES (SHEEP).//P50415
  - F-HEMBA1005737//CALCINEURIN B SUBUNIT (PROTEIN PHOSPHATASE 2B REGULATORY SUBUNIT),//
    4.5e-18:167:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P25296
- 45 F-HEMBA1005746
  - F-HEMBA1005755//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//7.4e-30:69:65//HOMO SAPIENS (HU-MAN) //P08547
  - F-HEMBA1005765//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.8e-19:60:63//HOMO SAPIENS (HU-MAN) //P39194
  - F-HEMBA1005780//METALLOTHIONEIN-I (MT-1).//1.0:31:38//COLUMBA LIVIA (DOMESTIC PIGEON).// P15786
    - F-HEMBA1005813
    - F-HEMBA1005815//CALPAIN, LARGE [CATALYTIC] SUBUNIT (EC 3.4.22.17) (CALCIUM- ACTIVATED NEU-TRAL PROTEINASE) (CANP) (MU/M-TYPE).//1.0e-23:200:31//GALLUS GALLUS (CHICKEN).//P00789
- 55 F-HEMBA1005822//PROTEIN Q300.//0.0016:21:80//MUS MUSCULUS (MOUSE).//Q02722 F-HEMBA1005829//III ALU SUBFAMILY SQ WARNING ENTRY !!!!//9.6e-33:96:73//HOMO SAPIENS (HUMAN).// P39194
  - F-HEMBA1005834//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG, I/1.6e-22:103:46//NYCTICEBUS COU-

CANG (SLOW LORIS) //P08548

F-HEMBA1005852//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//8.8e-06:95:35//MUS MUSCULUS (MOUSE).//P05143

F-HEMBA1005853//HYPOTHETICAL PROTEIN

MJ0647.//0.39:28:39//METHANOCOCCUS JANNASCHII.//Q58063

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F-HEMBA1005891//HYPOTHETICAL PROTEIN MTH137.//0.95:51:27//METHANOBACTERIUM THERMOAU-TOTROPHICLIM //026240

F-HEMBA1005894//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.6e-29:81:71//HOMO SAPIENS (HUMAN).//

P39195
F-HEMBA1005909/HYPOTHETICAL 8.2 KD PROTEIN B0353.1 IN CHROMOSOME III.//0.98:19:52//
CAENORHABDITIS ELEGANS.//010958

F-HEMBA1005911//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.9e-27:86:70//HOMO SAPIENS (HUMAN).// P39188

15 F-HEMBA1005921//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/1.3e-38:99:81//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1005931//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//2.3e-17:76:51//HOMO SAPI-ENS (HUMAN).//P51522

F-HEMBA1005934//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//0.024;54;40//HOMO SAPIENS (HUMAN).// P39189

F-HEMBA1005962

F-HEMBA 1005902

F-HEMBA1005963//COATOMER GAMMA SUBUNIT (GAMMA-COAT PROTEIN) (GAMMA-COP).//1.7e-32:89: 79//BOS TAURUS (BOVINE).//P53620

F-HEMBA1005990//HYPOTHETICAL BHLF1 PROTEIN.//3.0e-09:180:36//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181

F-HEMBA1005991//HYPOTHETICAL PROTEIN KIAA0032.//3.0e-17:107:43//HOMO SAPIENS (HUMAN).// Q15034

F-HEMBA1005999

F-HEMBA1006002

F-HEMBA1006005//CORNIFIN B (SMALL PROLINE-RICH PROTEIN 1B) (SPR1B) (SPR1 B).//0.0017:45:44// MUS MUSCULUS (MOUSE).//Q62267

F-HEMBA1006031//BASIC PROLINE-RICH PEPTIDE IB-1.//0.00016:84:39//HOMO SAPIENS (HUMAN).// P04281

F-HEMBA1006035//DNAK PROTEIN 1 (HEAT SHOCK PROTEIN 70) (HSP70).//0.43:100:27//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//Q55154

F-HEMBA1006036///!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//6.2e-64:150:74//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1006042

F-HEMBA1006067//METALLOTHIONEIN A (MT-A).//0.86:34:41//THERMARCES CERBERUS.//P52721 F-HEMBA1006081

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F-HEMBA1006090//SODIUM/GLUCOSE COTRANSPORTER 3 (NA(+)/GLUCOSE COTRANSPORTER 3) (LOW AFFINITY SODIUM-GLUCOSE COTRANSPORTER).//0.87:35:54//SUS SCROFA (PIG).//P31636

F-HEMBA1006091//EARLY NODULIN 20 PRECURSOR (N-20).//0.027:87:32//MEDICAGO TRUNCATULA (BAR-REL MEDIC).//P93329

45 F-HEMBA1006100//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//8.1e-09:58:60//HOMO SAPIENS (HUMAN).// P39195

F-HEMBA1006108/HYPOTHETICAL 56 6 KD PROTEIN IN URE2-SSU72 INTERGENIC REGION.//5.6e-16:88: 36/SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53867
F-HEMBA1006121/H/OMEOBOX PROTEIN CDX-1 (CAUDAL-TYPE HOMEOBOX PROTEIN 1).//3.4e-05:106:

F-HEMBA1006121//HOMEOBOX PROTEIN CDX-1 (CAUDAL-TYPE HOMEOBOX PROTEIN 1).//3.4e-05:106 37//HOMO SAPIENS (HUMAN).//P47902

F-HEMBA1006124//50S RIBOSOMAL PROTEIN L33.//1.0:12:83//BACILLUS STEAROTHERMOPHILUS.// P23375

F-HEMBA1006130//SEL-10 PROTEIN.//7.7e-05:129:28//CAENORHABDITIS ELEGANS.//Q93794

F-HEMBA1006138//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.8e-13:41:73//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBA1006142//!!!! ALU SUBFAMILY SC WARNING ENTRY!!!!//2.3e-39:101:77//HOMO SAPIENS (HU-MAN).//P39192

F-HEMBA1006155//GENE 33 POLYPEPTIDE.//0.21:70:31//RATTUS NORVEGICUS (RAT).//P05432

- F-HEMBA1006158
- F-HEMBA1006173//PROTEIN-TYROSINE PHOSPHATASE STRIATUM-ENRICHED (EC 3.1.3.48) (STEP) (NEU-RAL-SPECIFIC PROTEIN-TYROSINE PHOSPHATASE) (FRAGMENT)///0.017:20:95//HOMO SAPIENS (HU-MAN) //P54879
- F-HEMBA1006182///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.37:31:61//HOMO SAPIENS (HUMAN).//
  - F-HEMBA1006198//HOMEOBOX PROTEIN HOX-B3 (HOX-2.7) (MH-23).//0.85:61:29//MUS MUSCULUS (MOUSE).//P09026
    - F-HEMBA1006235//50S RIBOSOMAL PROTEIN L33.//1.0:26:38//AQUIFEX AEOLICUS.//O67756
- F-HEMBA1006248//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.0041:84:37//DROSOPHILA MELA-NOGASTER (FRUIT FLY) //001643
  - F-HEMBA1006252//BOWMAN-BIRK TYPE PROTEINASE INHIBITOR DE-3.//1.0:22:40//DOLICHOS AXILLARIS (MACROTYLOMA AXILLARE).//P01057
  - F-HEMBA1006253//DISINTEGRIN ERISTICOPHIN (PLATELET AGGREGATION ACTIVATION INHIBITOR).//
    0.95/19/47//ERISTOCOPHIS MACMAHONI (LEAF-NOSED VIPER).//P22826
- F-HEMBA1006259

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- F-HEMBA1006268//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//7.0e-05:32:65//HOMO SAPIENS (HUMAN).// P39192
- F-HEMBA1006272//RETROVIRUS-RELATED GAG POLYPROTEIN (VERSION 2).//4.8e-112:248:78//HOMO SA-PIENS (HUMAN).//P10264
- F-HEMBA1006278//POLY(A) POLYMERASE (EC 2.7.7.19) (PAP) (POLYNUCLEOTIDE ADENYLYLTRANS-FERASE) (FRAGMENT).//2.5e-71:164:75//HOMO SAPIENS (HUMAN).//P51003
  - F-HEMBA1006283//50S RIBOSOMAL PROTEIN L32.//0.81:27:44//THERMUS AQUATICUS (SUBSP. THER-MOPHILUS).//P80339
- 25 F-HEMBA1006284//CUTICLE COLLAGEN 2.//0.36:42:40//CAENORHABDITIS ELEGANS.//P17656
  - F-HEMBA1006291//HYPOTHETICAL 43.3 KD PROTEIN IN EVGS-GLK INTERGENIC REGION.//2.4e-37:143: 31//ESCHERICHIA COLL//P76518
  - F-HEMBA 1006293/MYELIN-OLIGODENDROCYTE GLYCOPROTEIN PRECURSOR //0.20:134:29//RATTUS NORVEGICUS (RAT) //0.63345 F-HEMBA 1006309//HYPOTHETICAL 54.2 KD PROTEIN IN ERP5-ORC6 INTERGENIC REGION //2.16-43:187-48/SACCHAROMYCES CEREVISIAE (BAKER'S YEASTI.//P38821
- F-HEMBA1006310//SIGNAL TRANSDUCER CD24 PRECURSOR (HEAT STABLE ANTIGEN) (HSA) (NECTADRIN).//0.71:46:39//RATTUS NORVEGICUS (RAT).//Q07490
  - F-HEMBA1006328//RNA POLYMERASE ALPHA SUBUNIT (EC 2.7.7.48) (NUCLEOCAPSID PHOSPHOPRO-TEIN).//0.44:141:24//HUMAN PARAINFLUENZA 1 VIRUS (STRAIN CI-5/73).//P32531
- F-HEMBA1006334//HYPOTHETICAL TRANSCRIPTIONAL REGULATOR AF1627.//0.98:26:46//ARCHAE-OGLOBUS FULGIDUS.//028646
  - F-HEMBA1006344//EZRIN (P81) (CYTOVILLIN) (VILLIN-2).//8.8e-08:91:36//MUS MUSCULUS (MOUSE).// P26640
- F-HEMBA1006347//MALES-ABSENT ON THE FIRST PROTEIN (EC 2.3.1.-).//9.1e-48:149:50//DROSOPHILA

  MELANOGASTER (FRUIT FLY).//02193
  - F-HEMBA1006349//METALLOTHIONEIN-LIKE PROTEIN 1.//0.015:59:33//CASUARINA GLAUCA (SWAMP OAK) //039511
  - F-HEMBA1006359//ZINC FINGER PROTEIN 43 (ZINC PROTEIN HTF6).//6.8e-96:261:66//HOMO SAPIENS (HU-MAN) //098460
- 45 F-HEMBA1006364//PUTATIVE ENDONUCLEASE C1F12.06C (EC 3.1.-.-).//0.97:60:35//SCHIZOSACCHARO-MYCES POMBE (FISSION YEAST).//O10348
  - F-HEMBA1006377//EARLY NODULIN 20 PRECURSOR (N-20).//0.00023:110:35//MEDICAGO TRUNCATULA (BARREL MEDIC).//P93329
    - F-HEMBA1006380
- 50 F-HEMBA1006381//METALLOTHIONEIN-II.//1.0:26:38//CANDIDA GLABRATA (YEAST) (TORULOPSIS GLA-BRATA).//P15114
  - F-HEMBA1006398//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.3e-26:123:52//HOMO SAPIENS (HU-MAN).//P08547
  - F-HEMBA1006416
- 55 F-HEMBA1006419//!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.2e-24:102:50//HOMO SAPIENS (HU-MAN) //P39189
  - F-HEMBA1006421///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.1e-21:101:57//HOMO SAPIENS (HUMAN).// P39188

- F-HEMBA1006424//HYPOTHETICAL PROTEIN IORF1.//0.85:55:30//BOVINE CORONAVIRUS (STRAIN MEBUS), AND BOVINE CORONAVIRUS (STRAIN QUEBEC).//P22053
- F-HEMBA1006426//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.8e-36:78:74//HOMO SAPIENS (HUMAN).//
- F-HEMBA1006438//HYPOTHETICAL 8.1 KD PROTEIN (ORF65).//1.0:38:36//GUILLARDIA THETA (CRYPTO-MONAS PHI).//O78421
  - F-HEMBA1006445//RAS-LIKE PROTEIN 3.//1.9e-06:40:47//RHIZOMUCOR RACEMOSUS (MUCOR CIRCINEL-LOIDES F. LUSITANICUS) //P22280
  - F-HEMBA1006446

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- F-HEMBA1006461/!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//4.1e-18:68:67//HOMO SAPIENS (HUMAN).// P39192
  - F-HEMBA1006467
  - F-HEMBA1006471
  - F-HEMBA1006474//40 KD PROTEIN.//1.1e-37:231:38//BORNA DISEASE VIRUS (BDV).//Q01552
- 15 F-HEMBA1006483//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//6.1e-38:77:74//HOMO SAPIENS (HUMAN).// P39192
  - F-HEMBA1006485//HYPOTHETICAL 9.3 KD PROTEIN IN NAD3-NAD7 INTERGENIC REGION (ORF 79).//0.91: 30:40//MARCHANTIA POLYMORPHA (LIVERWORT).//P38465
  - F-HEMBA1006486//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.1e-12:78:51//HOMO SAPIENS (HU-MAN) //P08547
  - F-HEMBA1006489//FUN34 PROTEIN://0.94:58:37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
    - F-HEMBA1006492//NADH-UBIQUINONE OXIDOREDUCTASE MWFE SUBUNIT (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-MWFE) (CI-MWFE).//0.87:44:36//HOMO SAPIENS (HUMAN).//O15239
- F-HEMBA1008494//FERREDOXIN-LIKE PROTEIN IN NIF REGION.//0.11:46:26//RHIZOBIUM LEGUMINOSA-RUM (BIOVAR TRIFOLII).//P42711 F-HEMBA1006402
  - F-HEMBA1006502//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.15:26:73//HOMO SAPIENS (HUMAN).//
- F-HEMBA1006507//DIAPHANOUS PROTEIN.//0.0055:129:28//DROSOPHILA MELANOGASTER (FRUIT FLY).// PARROR
  - F-HEMBA1006521//3-OXOACYL-[ACYL-CARRIER PROTEIN] REDUCTASE (EC 1.1.1.100) (3-KETOACYL-ACYL CARRIER PROTEIN REDUCTASE)//1.1e-32:177:41//ESCHERICHIA COLI.//P25716
    - F-HEMBA1006530//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3) (FRAGMENT).//0.052:84: 26//LEISHMANIA TARENTOLAE (SAUROLEISHMANIA TARENTOLAE).//P15583
      - F-HEMBA1006535//INHIBITOR OF APOPTOSIS PROTEIN 1 (MIAP1) (MIAP-1).//6.6e-05:53:39//MUS MUSCU-LUS (MOUSE).//O08863
      - F-HEMBA1006540//PRESYNAPTIC PROTEIN SAP97 (SYNAPSE-ASSOCIATED PROTEIN 97) (DISCS, LARGE HOMOLOG 1).//2.1e-07:206:23//RATTUS NORVEGICUS (RAT).//Q62696
- 40 F-HEMBA1006546//PROBABLE ES PROTEIN //0 .11:70.32//HUMAN PAPILLOMAVIRUS TYPE 51//P26553 F-HEMBA1006559//SUPPRESSOR PROTEIN SRP40.//0.015:221.20//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEASTI./P32583
  - F-HEMBA1006562//SALIVARY PROLINE-RICH PROTEIN PO PRECURSOR (ALLELE S).//1.5e-07:122:33//HO-MO SAPIENS (HUMAN).//P10163
- 45 F-HEMBA106866/CELL DIVISION PROTEIN KINASE 2 (EC 2.7.1.-) (CDC2 HOMOLOG EG1 PROTEIN KINASE)/I0.63:53:37/IXENOPUS LAEVIS (AFRICAN CLAWED FROG).//P23437
  - F-HEMBA1006569//COLLAGEN ALPHA 2(I) CHAIN (FRAGMENT).//4.4e-06:88:39//BOS TAURUS (BOVINE).// P02465
    - F-HEMBA1006579
- F-HEMBA1006583//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//0.011:61:40//MUS MUSCULUS (MOUSE).//P05142
  - F-HEMBA1006595///III ALU SUBFAMILY SQ WARNING ENTRY IIII//5.6e-34:93:77//HOMO SAPIENS (HU-MAN).//P39194
- F-HEMBA1006597//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.9e-26:75:74//HOMO SAPIENS (HUMAN).//
  P30105
- F-HEMBA1006612//SUPPRESSOR PROTEIN SRP40.//0.026:221:22//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583
  - F-HEMBA1006617/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/6.6e-20:73:63//HOMO SAPIENS (HUMAN).//

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F-HEMBA1006624//HYPOTHETICAL 41.9 KD PROTEIN IN SDS3-THS1 INTERGENIC REGION.//2.6e-31:209: 44//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40506

F-HEMBA1006631//HYPOTHETICAL 62.8 KD PROTEIN IN TAF145-YOR1 INTERGENIC REGION.//1.5e-15: 131:41//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53331

### F-HEMBA1006635

F-HEMBA1006639//POLYADENYLATE-BINDING PROTEIN 1 (POLY(A) BINDING PROTEIN 1) (PABP 1).//2.2e-11:48:75//MUS MUSCULUS (MOUSE).//P29341

F-HEMBA1006643//LONG NEUROTOXIN CR1 PRECURSOR (KAPPA NEUROTOXIN).//0.28:48:27//BUNGA-RUS MULTICINCTUS (MANY-BANDED KRAIT).//P15817

F-HEMBA1006648//ZINC FINGER PROTEIN 12 (ZINC FINGER PROTEIN KOX3) (FRAGMENT),//0.26:17:47// HOMO SAPIENS (HUMAN),//P17014

F-HEMBA1006652//60S RIBOSOMAL PROTEIN L7.//2.4e-44:206:47//MUS MUSCULUS (MOUSE).//P14148

# F-HEMBA1006653

15 F-HEMBA 1006659

F-HEMBA1006665//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.018:43:58//HOMO SAPIENS (HU-

F-HEMBA1006674//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII135) (TAFII-130) (TAFII130).//2.9e-05:154:33//HOMO SAPIENS (HUMAN).//O00268

20 F-HEMBA1006676//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//3.6e-09:52:51//OWENIA FUSI-FORMIS //P21260

### F-HEMBA1006682

F-HEMBA1006695//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/2.1e-06:35:65//HOMO SAPIENS (HUMAN).// P39188

### 25 F-HEMBA1006696

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F-HEMBA1006708//HYPOTHETICAL 46.4 KD TRP-ASP REPEATS CONTAINING PROTEIN IN PMC1-TFG2 INTERGENIC REGION.//3.4e-19:104:45//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53196

F-HEMBA1006709//RETINOIC ACID RECEPTOR RXR-BETA.//0.24:111:36//HOMO SAPIENS (HUMAN).// P28702

### 30 F-HEMBA1006717

F-HEMBA1006737//ANKYRIN, BRAIN VARIANT 2 (ANKYRIN B) (ANKYRIN, NONERYTHROID) (FRAGMENT).// 5.8e-09:111:40//HOMO SAPIENS (HUMAN).//Q01485

F-HEMBA1006744//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//1.8e-32:84:78//HOMO SAPIENS (HU-MAN).//P39191

35 F-HEMBA1006754//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.3e-75:220:62//HOMO SAPIENS (HU-MAN),//P08547

F-HEMBA1006758//VASCULAR ENDOTHELIAL-CADHERIN PRECURSOR (VECADHERIN) (CADHERIN-5) (7B4 ANTIGEN) (CD144 ANTIGEN).//0.024:110:29//HOMO SAPIENS (HUMAN).//P33151 F-HEMBA 1006767

F-HEMBA1006779//MITOCHONDRIAL RIBOSOMAL PROTEIN \$12://0.67:19:42//LEISHMANIA TARENTOLAE 40 (SAUROLEISHMANIA TARENTOLAE).//Q34940 F-HEMBA1006780

F-HEMBA1006789//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.056:98:30//MUS MUSCULUS (MOUSE) //P05143

45 F-HEMBA1006795//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//2.9e-11:143:30//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548

F-HEMBA1006796//WISKOTT-ALDRICH SYNDROME PROTEIN HOMOLOG (WASP).//0.16:38:42//MUS MUS-CULUS (MOUSE),//P70315

F-HEMBA1006807//HYPOTHETICAL 46.4 KD PROTEIN T16H12.5 IN CHROMOSOME III.//4.4e-75:184:77// CAENORHABDITIS ELEGANS.//P34568

F-HEMBA1006821//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//0.011:20:85//HOMO SAPIENS (HUMAN).// P3Q1Q4

F-HEMBA1006824//PROTEIN B11.//0.44:27:44//VACCINIA VIRUS (STRAIN WR).//Q01229

F-HEMBA1006832//HYPOTHETICAL 34.6 KD PROTEIN C13G5.2 IN CHROMOSOME III.//1.0:46:36// CAENORHABDITIS ELEGANS.//P34327

### F-HFMBA1006849

F-HEMBA1006865//ACROSIN INHIBITORS IIA AND IIB (BUSI-II).//1.0:41:31//BOS TAURUS (BOVINE).//P01001 F-HEMBA1006877//OXYSTEROL-BINDING PROTEIN.//3.7e-26:239:36//ORYCTOLAGUS CUNICULUS (RAB-

BIT).//P16258

F-HEMBA1006885//HYPOTHETICAL 27.2 KD PROTEIN F09E5.8 IN CHROMOSOME II.//4.5e-38:185:43//

F-HEMBA1006900

F-HEMBA1006914//UBIQUITIN-ACTIVATING ENZYME E1-LIKE (POLYMERASE-INTERACTING PROTEIN 2).//
5.2e-27:269:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P52488

F-HEMBA1006921//CYTOTOXIN 3 (COMPONENT 3.20).//0.99:32:37//NAJA MELANOLEUCA (FOREST CO-BRA) (BLACK-LIPPED COBRA).//P01473

F-HEMBA1006926//ATROPHIN-1 (DENTATORUBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN).//0.0024:148: 33//RATTUS NORVEGICUS (RAT).//P54258

F-HEMBA1006929/HYPOTHETICAL PROTEIN MJ0525.//0.95:35:20//METHANOCOCCUS JANNASCHII.//

F-HEMBA1006936//SALIVARY ACIDIC PROLINE-RICH PHOSPHOPROTEIN 1/2 PRECURSOR (PRP-1 / PRP-3) (PRP-2) (PRP-4) (PIF-F) PIF-2) (PROTEIN A / PROTEIN C) (CONTAINS: PEPTIDE P-C].//0.074:116:31//HOMO SAPIENS (HUMAN)./P02810

15 SAPIENS (HUMAN F-HEMBA1006938

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F-HEMBA1006941/THIOREDOXIN H-TYPE 1 (TRX-H1).//2.1e-13:90:33//NICOTIANA TABACUM (COMMON TOBACCO).//P29449

F-HEMBA1006949

20 F-HEMBA1006973//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.75:29:55//BOS TAURUS (BOVINE).// P25508

P25508 F-HEMBA1006976//CMP-N-ACETYLNEURAMINATE-BETA-GALACTOSAMIDE-ALPHA-2,3-SIALYLTRANS-FERASE (EC 2 4 99 - 1/BETA-GALACTOSIDE ALPHA-2-3-SIALYLTRANSFERASE) (ST3GALIII) (ALPHA 2.3-ST)

(GAL-NAC6S) (STZ) (SIAT4-C) (SAT-3) (ST-4).//3.9e-108:117:95//HOMO SAPIENS (HUMAN).//Q11206 F-HEMBA1006993

F-HEMBA1006996//HYPOTHETICAL 8.7 KD PROTEIN IN RPL22-RPL23 INTERGENIC REGION (ORF70).//
0.12:51:33//ASTASIA LONGA (EUGLENOPHYCEAN ALGA).//P34779

F-HEMBA1007002//PLATELET GLYCOPROTEIN IX PRECURSOR (GPIX) (CD42A).//0.00096:60:33//HOMO SA-PIENS (HUMAN).//P14770

39 F-HEMBA1007017//HYPOTHETICAL 7.2 KD PROTEIN IN CYAY-DAPF INTERGENIC REGION.//1.0:25:56//ES-CHERICHIA COLI.//P39166

F-HEMBA1007018//DYNEIN LIGHT INTERMEDIATE CHAIN 1, CYTOSOLIC (LIC57/59) (DYNEIN LIGHT CHAIN A) (DLC-A).//8.5e-120:278:80//GALLUS GALLUS (CHICKEN).//Q90828

F-HEMBA1007045//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//2.1e-12:158:29// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437

F-HEMBA1007051

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F-HEMBA1007052//60S RIBOSOMAL PROTEIN L37-B (L35) (YP55).//0.94:37:35//SACCHAROMYCES CERE-VISIAE (BAKER'S YEAST).//P51402

F-HEMBA1007062//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.93:55:29//RHINOCEROS UNICORNIS (GREATER INDIAN RHINOCEROS).//096063

F-HEMBA1007066//ECLOSION HORMONE PRECURSOR (ECDYSIS ACTIVATOR) (EH).//0.58:49:38//BOM-BYX MORI (SILK MOTH).//P25331

F-HEMBA1007073//PUTATTVE SMALL MEMBRANE PROTEIN (ORF 4).//0.86:46:34//CANINE ENTERIC CORONAVIRUS (STRAIN INSAVC-1) (CCV).//P36696

45 F-HEMBA1007078//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//8.6e-29:56:67//HOMO SAPIENS (HUMAN).// P39193

F-HEMBA1007080//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//0.028:122:30//XENO-PUS LAEVIS (AFRICAN CLAWED FROG).//P17437

F-HEMBA1007085//RTOA PROTEIN (RATIO-A).//7.4e-11:221:31//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P54681

F-HEMBA1007087//HYPOTHETICAL PROTEIN MJ0162.//3.3e-29:173:36//METHANOCOCCUS JANNASCHII.// Q57626

F-HEMBA1007112

F-HEMBA1007113

55 F-HEMBA1007121//INOSITOL POLYPHOSPHATE 1-PHOSPHATASE (EC 3.1.3.57) (IPP).//5.4e-07:90:28//HO-MO SAPIENS (HUMAN).//P49441

F-HEMBA1007129//HIRUSTASIN.//0.88:37:32//HIRUDO MEDICINALIS (MEDICINAL LEECH) .//P80302 F-HEMBA1007147//HYPOTHETICAL 12.0 KD PROTEIN IN DST1-HEM2 INTERGENIC REGION.//0.92:23:34//

- SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P53182
- F-HEMBA1007149//BACTERIOCIN MICROCIN B17 PRECURSOR (MCB17).//0.0078:17:70//ESCHERICHIA
- F-HEMBA1007151//WDNM1 PROTEIN PRECURSOR.//0.25:45:37//MUS MUSCULUS (MOUSE).//Q62477
- F-HEMBA1007174//HYPOTHETICAL 45.1 KD PROTEIN IN RPS5-ZMS1 INTERGENIC REGION.//6.9e-18:97: 47//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST\.//P47160
  - F-HEMBA1007178//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/9.8e-06:38:65//HOMO SAPIENS (HUMAN).// P39195
- F-HEMBA1007194//GLUCOSE-6-PHOSPHATE 1-DEHYDROGENASE, CHOLOROPLAST ISOFORM PRECUR-SOR (EC. 1.1.1.49) (G6PD).//1.0:80:32//NICOTIANA TABACUM (COMMON TOBACCO).//Q43793
- F-HEMBA1007203//PROTEIN A22.//1.0:115:26//VARIOLA VIRUS.//P33845
  - F-HEMBA1007206
  - F-HEMBA1007224//HYPOTHETICAL 35.7 KD PROTEIN C41C4.6 IN CHROMOSOME II.//2.4e-05:92:30// CAENORHARDITIS FLEGANS //009275
- 15 F-HEMBA1007243/HYPOXANTHINE-GUANINE PHOSPHORIBOSYLTRANSFERASE (EC 2.4.2.8) (HGPRT) (HGPRTASE) (HPRT B).//3.1e-74:205:67//MUS MUSCULUS (MOUSE).//P00493
  - F-HEMBA1007251/MTELLINE MEMBRANE PROTEIN VM26AB PRECURSOR (PROTEIN TU-4) (PROTEIN SV23)://0.52:108:30//DROSOPHILA MELANOGASTER (FRUIT FLY)://P13238
    E-HEMBA1007256
  - F-HEMBA1007267//CALICIN (FRAGMENT).//0.060:88:31//HOMO SAPIENS (HUMAN).//Q13939
    - F-HEMBA1007273//HYPOTHETICAL 8.1 KD PROTEIN (ORF65).//0.95:40:37//GUILLARDIA THETA (CRYPTO-MONAS PHI).//078421
    - F-HEMBA1007279//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.6e-24:98:64//HOMO SAPIENS (HUMAN).//
- 25 F-HEMBA1007281

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- F-HEMBA1007288//HYPOTHETICAL 13.5 KD PROTEIN IN ZMS1-MNS1 INTERGENIC REGION.//0.88:11:54// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47162
- F-HEMBA1007300//CGMP-SPECIFIC 3',5'-CYCLIC PHOSPHODIESTERASE (EC 3.1.4.17) (CGB-PDE).//2.7e-43:220:41//BOS TAURUS (BOVINE).//Q28156
- 30 F-HEMBA107301//PROCOLLAGEN ALPHA 1(III) CHAIN PRECURSOR.//3.36-22:115:33//HOMO SAPIENS (HUMAN),//P02461 F-HEMBA107319
  - F-HEMBA1007320//HYPOTHETICAL 28.0 KD PROTEIN IN GLOB-RNHA INTERGENIC REGION.//1.0:48:37// ESCHERICHIA COLI.//P75672
- F-HEMBA1007322/THREONINE DEHYDRATASE OPERON ACTIVATOR PROTEIN.//1.0:59:33//ESCHERICHIA COLI.//P11866
  - F-HEMBA1007327
  - F-HEMBA1007341//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.1e-12:37:62//HOMO SAPIENS (HUMAN).//
- 40 F-HEMBA1007342//PROBABLE E5 PROTEIN.//0.89:96:29//PYGMY CHIMPANZEE PAPILLOMAVIRUS TYPE 1 //O02268
  - F-HEMBA1007347//INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN 2 PRECURSOR (IGFBP-2) (IBP-2) (IGF-BINDING PROTEIN 2).//0.92:62:43//OVIS ARIES (SHEEP).//Q29400
  - F-HEMBB1000005//WEAK NEUROTOXIN 5.//0.98:30:33//NAJA NAJA (INDIAN COBRA).//P29179
- 45 F-HEMBB1000008//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/2.7e-35:73:84//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBB1000018/HYPOTHETICAL BHLF1 PROTEIN./0.39:90:37//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4)./P03181 F-HEMBB1000024/VIRE LOCUS 9 KD VIRULENCE PROTEIN.//0.66:36-41//AGROBACTERIUM TUMEFA-
  - F-HEMBB1000024//VIRE LOCUS 9 KD VIRULENCE PROTEIN.//0.66:36:41//AGROBACTERIUM TUMEFA-CIENS.//P08061
- F-HEMBB1000025//MUSCARINIC TOXIN ALPHA (MT-ALPHA).//0.46:32:40//DENDROASPIS POLYLEPIS POLYLEPIS (BLACK MAMBA).//P80494
  - F-HEMBB1000030//SUPPRESSOR PROTEIN SRP40.//6.7e-07:50:52//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32583
- 55 F-HEMBB1000036//HYPOTHETICAL 43.2 KD PROTEIN C34E10.1 IN CHROMOSOME III.//2.5e-07:120:29//
  CAFNORHABDITIS FLEGANS //P46576
  - F-HEMBB1000037//HYPOTHETICAL 59.9 KD PROTEIN-IN SGA1-KTR7 INTERGENIC REGION.//1.7e-05:71: 29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40492

- F-HEMBB1000039//VERY HYPOTHETICAL 11.9 KD PROTEIN C4H3.12C IN CHROMOSOME I.//1.0:61:21// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10219
- F-HEMBB1000044

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- F-HEMBB1000048//HYPOTHETICAL 15.7 KD PROTEIN IN IDH-DEOR INTERGENIC REGION.//1.0:63:31//BA-CILLUS SUBTILIS //P54942
- F-HEMBB1000050//!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//9.0e-14:34:79//HOMO SAPIENS (HU-MAN) //P30194
  - F-HEMBB1000054//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//5.9e-31:45:73//HOMO SAPIENS (HUMAN).// P39193
- F-HEMBB1000055//MUSCARINIC TOXIN ALPHA (MT-ALPHA).//1.0:14:57//DENDROASPIS POLYLEPIS POLYLEPIS (BLACK MAMBA).//P80494
  - F-HEMBB1000059///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.0e-21:82:59//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBB1000083//CHROMOGRANIN A PRECURSOR (CGA) [CONTAINS: PANCREASTATIN; BETA-GRANIN; WE-14].//0.87:172:28//RATTUS NORVEGICUS (RAT).//P10354
  - F-HEMBB1000089//HYPOTHETICAL 9.5 KD PROTEIN IN SPEA-METK INTERGENIC REGION (F83).//1.0:42: 33//ESCHERICHIA COLI.//P46879
    - F-HEMBB1000099//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//7.7e-08:31:87//HOMO SAPIENS (HUMAN).//
- F-HEMBB1000103/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.4e-38:136:58//HOMO SAPIENS (HU-MAN).//P08547
- F-HEMBB1000119//MAF PROTEIN://3.6e-32:195:43//BACILLUS SUBTILIS://Q02169
- 25 F-HEMBB1000136//HYPOTHETICAL 12.7 KD PROTEIN IN PCS60-ABD1 INTERGENIC REGION.//0.65:71:32// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38327
  - F-HEMBB1000141//B-CELL GROWTH FACTOR PRECURSOR (BCGF-12 KD).//0.00014:34:64//HOMO SAPI-ENS (HUMAN).//P20931
  - F-HEMBB1000144//!!!! ALU SUBFAMILY SB2 WARNING ENTRY!!!!//2.0e-26:81:69//HOMO SAPIENS (HU-MAN).//P39191
- F-HEMBB1000173//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.2e-29:91:71//HOMO SAPIENS (HUMAN).//
  - F-HEMBB1000175//ANTIMICROBIAL PEPTIDE ENAP-1 (FRAGMENT).//0.97:41:36//EQUUS CABALLUS (HORSE).//P80930
- 35 F-HEMBB1000198//HYPOTHETICAL 7.7 KD PROTEIN YCF33 (ORF67).//0.91:21:52//PORPHYRA PURPUREA.//P51329
  - F-HEMBB1000215///!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//3.4e-08:39:76//HOMO SAPIENS (HUMAN).// P39192
- F-HEMBB1000217//DNA DAMAGE TOLERANCE PROTEIN RHC31 (RAD31 HOMOLOG).//2.9e-32:174:40//
  SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//O06624
  - F-HEMBB1000218//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.73:31:38//MICROTUS PENNSYLVAN-ICUS (MEADOW VOLE).//P24949
    - F-HEMBB1000226//HYPOTHETICAL 37.0 KD PROTEIN B0495.8 IN CHROMOSOME II.//6.5e-26:191:34//
      CAENORHARDITIS FLEGANS //Q09217
- 45 F-HEMBB1000240
  - F-HEMBB1000244//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.9e-05:44:61//HOMO SAPIENS (HUMAN).// P39188
    - F-HEMBB1000250
  - F-HEMBB1000258
- F-HEMBB1000264//CUTICLE COLLAGEN SQT-1.//0.15:89:33//CAENORHABDITIS ELEGANS.//P12114
  - F-HEMBB1000266//TRANSLATION INITIATION FACTOR IF-2.//2.7e-06:167:22//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P39730
  - F-HEMBB1000272//CYTOCHROME C OXIDASE POLYPEPTIDE VIB (EC 1.9.3.1) (AED).//0.75:30:43//BOS TAU-RUS (BOVINE).//P00429
- 55 F.HEMBB1000274/ICORNIFIN (SMALL PROLINE-RICH PROTEIN I) (SPR-I) (SMALL PROLINE-RICH SQUA-MOUS CELL MARKER) (SPRP)./I/1.0:38:36//SUS SCROFA (PIG)./I/P35323
  - F-HEMBB1000284//CALTRIN (CALCIUM TRANSPORT INHIBITOR).//1.0:56:30//MUS MUSCULUS (MOUSE).// Q09098

- F-HEMBB1000307
- F-HEMBB1000312

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- F-HEMBB1000317//THROMBOSPONDIN 1 PRECURSOR.//3.2e-32:135:43//HOMO SAPIENS (HUMAN).// PO7006
- F-HEMBB1000318//PUTATIVE SMALL MEMBRANE PROTEIN (NONSTRUCTURAL PROTEIN NS3) (NON-STRUCTURAL 9.5 KD PROTEIN).//0.41:51:31//HUMAN CORONAVIRUS (STRAIN OC43).//Q04854
  - F-HEMBB1000335//ZINC FINGER PROTEIN 13 (ZFP-13) (KROX-8 PROTEIN) (FRAGMENT).//0.82:33:45//MUS MUSCULUS (MOUSE).//P10754
  - F-HEMBB1000336//ALDEHYDE OXIDASE (EC 1.2.3.1) (FRAGMENTS).//0.80:44:40//ORYCTOLAGUS CUNIC-ULUS (RABBIT).//P80456
    - F-HEMBB1000337//SPLICING FACTOR, ARGININE/SERINE-RICH 4 (PRE-MRNA SPLICING FACTOR SRP75),//0.94:118:22//HOMO SAPIENS (HUMAN),//Q08170
  - F-HEMBB1000338//MALE SPECIFIC SPERM PROTEIN MST84DA.//0.042:33:39//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01642
- F-HEMBB1000339//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.2e-14:54:55//HOMO SAPIENS (HUMAN).// 15 P30188
  - F-HEMBB1000341//GENE 74 PROTEIN (GP74).//1.0:39:33//MYCOBACTERIOPHAGE L5.//Q05289
    - F-HEMBB1000343
  - F-HEMBB1000354///III ALU SUBFAMILY J WARNING ENTRY IIII//1.1e-15:83:56//HOMO SAPIENS (HUMAN).//
  - F-HEMBB1000369//PROTEIN Q300.//0.99:27:40//MUS MUSCULUS (MOUSE).//Q02722
    - F-HEMBB1000374//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!/4.7e-34:56:78//HOMO SAPIENS (HUMAN).//
    - F-HEMBB1000376
- 25 F-HEMBB1000391//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.0013:79:35//BOS TAURUS (BO-VINE) //P25508
  - F-HEMBB1000399//CHECKPOINT PROTEIN RAD17.//2.8e-15:187:31//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P50531
- F-HEMBB1000402//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3) (FRAGMENT),//0.027:60: 30 38//LEISHMANIA TARENTOLAE (SAUROLEISHMANIA TARENTOLAE).//P15583
  - F-HEMBB1000404//CYANELLE 50S RIBOSOMAL PROTEIN L28.//0.94:29:27//CYANOPHORA PARADOXA.// P48129
    - F-HEMBB1000420//SPLICEOSOME ASSOCIATED PROTEIN 49 (SAP 49) (SF3B53).//0.023;97:35//HOMO SA-PIENS (HUMAN).//Q15427
- 35 F-HEMBB1000434//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//4.8e-20:111:54//HOMO SAPIENS (HU-MAN).//P39194
  - F-HEMBB1000438//HYPOTHETICAL 7.9 KD PROTEIN IN GP55-NRDG INTERGENIC REGION://0.93:24:50// BACTERIOPHAGE T4 //P07076
- F-HEMBB1000441/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/4.4e-23:85:70//HOMO SAPIENS (HUMAN).// 40 P30188
  - F-HEMBB1000449/!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/0.88:27:51/HOMO SAPIENS (HUMAN).// P39195
    - F-HEMBB1000455
    - F-HEMBB1000472
- 45 F-HEMBB1000480//PROTEIN STBC.//1.0:52:30//ESCHERICHIA COLI.//P11905
  - F-HEMBB1000487//SHORT NEUROTOXIN 1 (NEUROTOXIN ALPHA) (NEUROTOXIN II).//0.93:29:34//NAJA OXIANA (CENTRAL ASIAN COBRA) (OXUS COBRA).//P01427
    - F-HEMBB1000490//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.3e-16:50:80//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBB1000491

- F-HEMBB1000493//3A PROTEIN.//1.0:51:35//AVIAN INFECTIOUS BRONCHITIS VIRUS (STRAIN BEAU-DETTE) (IBV).//P30237
- F-HEMBB1000510//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//9.7e-27:132:45//HOMO SAPIENS (HU-
- 55 F-HEMBB1000518//CYTOCHROME C OXIDASE POLYPEPTIDE III (EC 1.9.3.1).//0.021:47:40//LEISHMANIA TARENTOLAE (SAUROLEISHMANIA TARENTOLAE),//P14546
  - F-HEMBB1000523
  - F-HEMBB1000530//COLLAGEN ALPHA 1(XIV) CHAIN PRECURSOR (UNDULIN),//9.8e-14:43:83//GALLUS

GALLUS (CHICKEN) //P32018

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- F-HEMBB1000550/NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3).//0.19:97:30//TRYPANO-SOMA BRUCEI BRUCEI.//P04540
- F-HEMBB1000554//MATERNAL B9.10 PROTEIN (P30 B9.10).//0.94:82:25//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P40744
- F-HEMBB1000556//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII135) (TAFII-130) (TAFII130),//0.043:201:29//HOMO SAPIENS (HUMAN),//000268
- F-HEMBB1000564//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:5:2:34//METRIDIUM SENILE (BROWN SEA ANEMONE) (FRILLED SEA ANEMONE).//O47493
- F-HEMBB1000573//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//2.3e-10:52:73//HOMO SAPIENS (HU-MAN).//P39191
  - F-HEMBB1000575//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.8e-26:76:76//HOMO SAPIENS (HUMAN).// P39192
  - F-HEMBB1000586/NADH-UBIQUINONE OXIDOREDUCTASE MLRQ SUBUNIT (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-MLRQ) (CI-MLRQ) //0.74:23:52/HOMO SAPIENS (HUMAN) //000483
  - F-HEMBB1000589//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.9e-25:61:75//HOMO SAPIENS (HUMAN).//
    - F-HEMBB1000591/ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:34:35//PETROMYZON MARINUS (SEA LAMPREY).//035537
- F.-HEMBB1000592//SMALL PROLINE-RICH PROTEIN 2-1.//0.0016-49-42//HOMO SAPIENS (HUMAN).//P35326 F.-HEMBB1000593//COLLAGEN ALPHA 1(III) CHAIN (FRAGMENTS).//0.0070:189:32//GALLUS GALLUS (CHICKEN)./P2130
  - F-HEMBB1000598/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//5.7e-10:110:41//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548
- F-HEMBB1000623//HYPOTHETICAL 54.9 KD PROTEIN C02F5.7 IN CHROMOSOME III.//0.0022:98:28// CAENORHABDITIS ELEGANS.//P34284 E-JEMBR1000830
  - F-HEMBB1000631//ALPHA-2C-1 ADRENERGIC RECEPTOR (ALPHA-2C-1 ADRENOCEPTOR) (SUBTYPE C4).//8.8e-06:59:40//HOMO SAPIENS (HUMAN).//P18825
- F-HEMBB1000632//GUANINE NUCLEOTIDE RELEASING PROTEIN (GNRP).//7.3e-13:173:28//MUS MUSCU-LUS (MOUSE).//P27671
  - F-HEMBB1000637//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//4.6e-41:94:82//HOMO SAPIENS (HUMAN).// P39193
  - F-HEMBB1000638//INVOLUCRIN.//1.9e-06:144:29//HOMO SAPIENS (HUMAN).//P07476
- 35 F-HEMBB1000643//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//8.3e-30:77:76//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBB1000649//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.5e-37:58:81//HOMO SAPIENS (HUMAN).//
- F-HEMBB1000652//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.2e-37:61:77//HOMO SAPIENS (HUMAN).//
- F-HEMBB1000665//HYPOTHETICAL PROTEIN BBD24.//0.83:38:36//BORRELIA BURGDORFERI (LYME DIS-EASE SPIROCHETE).//P70845
  - F-HEMBB1000671//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.8e-51:74:71//HOMO SAPIENS (HU-
- 45 F-HEMBB1000673//HEAT-STABLE ENTEROTOXIN A3/A4 PRECURSOR (STA3/STA4) (ST-IB) (ST-II).//0.012: 37:37//ESCHERICHIA COLL.//P07965
  - F-HEMBB1000684//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//3.1e-21:66:72//HOMO SAPIENS (HUMAN).// P39193
    - F-HEMBB1000693//HUNTINGTIN ASSOCIATED PROTEIN 1 (HAP1).//5.2e-26:121:49//RATTUS NORVEGICUS (RAT).//P54256
    - F-HEMBB1000705

- F-HEMBB1000706
- F-HEMBB1000709//HYPOTHETICAL 5.8 KD PROTEIN.//1.0:29:44//CLOVER YELLOW MOSAIC VIRUS (CYMV).//P16485
- 55 F-HEMBB1000725//RAS-RELATED PROTEIN RAB-8B.//7.4e-105:205:98//RATTUS NORVEGICUS (RAT).//
  - F-HEMBB1000726//IIII ALU SUBFAMILY SQ WARNING ENTRY IIII//1.4e-25:85:70//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBB1000738//50S RIBOSOMAL PROTEIN L33.//1.0:41:31//THERMUS AQUATICUS (SUBSP. THER-MOPHILUS) //P35871

F-HEMBB1000749//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.3e-29:42:85//HOMO SAPIENS (HU-MAN) //P39194

F-HEMBB1000763//NIFU PROTEIN.//0.089:63:36//FRANKIA ALNI.//P46045

F-HEMBB100770//CALTRIN-LIKE PROTEIN II.//0.98:13:69//CAVIA PORCELLUS (GUINEA PIG).//P22075 F-HEMBB1000774//HIGH MOBILITY GROUP PROTEIN HMG-Y.//0.029:53:32//MUS MUSCULUS (MOUSE).// P17085

F-HEMBB1000781//MAPK/ERK KINASE KINASE 2 (EC 2.7.1.-) (MEK KINASE 2) (MEKK 2).//3.5e-75:144:98// MUS MUSCULUS (MOUSE) //061083

F-HEMBB1000789//PUTATIVE 90.2 KD ZINC FINGER PROTEIN IN CCA1-ADK2 INTERGENIC REGION.//2.6e-49:232-43//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39956

F-HEMBB1000790//!!!! ALU SUBFAMILY J WARNING ENTRY !!!//2.9e-16:93:51//HOMO SAPIENS (HUMAN).//

15 F-HEMBB1000794

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F-HEMBB1000807//MUSCARINIC ACETYLCHOLINE RECEPTOR M3.//0.54:111:27//GALLUS GALLUS (CHICK-

F-HEMBB1000810

F-HEMBB1000821

F-HEMBB1000822//HYPOTHETICAL 10 KD PROTEIN (ORF 6).//0.10:50:34//NARCISSUS MOSAIC VIRUS (NMM) //P15099

F-HEMBB1000826//B-CELL GROWTH FACTOR PRECURSOR (BCGF-12 KD).//0.00025:73:39//HOMO SAPIENS (HUMAN).//P20931

F-HEMBB1000827//HYPOTHETICAL 7.4 KD PROTEIN.//0.89:23:52//THERMOPROTEUS TENAX VIRUS 1

(STRAIN KRA1) (TTV1).//P19302 F-HEMBB1000831//MALE SPECIFIC SPERM PROTEIN MST87F.//0.98:35:40//DROSOPHILA MELA-

NOGASTER (FRUIT FLY).//P08175
F-HEMBE1000835/fL/INE-1 REVERSE TRANSCRIPTASE HOMOLOG///7.8e-31:96:46//HOMO SAPIENS (HU-MAN) //P08547

30 F-HEMBB100840//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.00012:102:36//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548

F-HEMBB1000848//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//7.3e-97:239:70//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBB1000852

F-HEMBB1000870

F-HEMBB1000876//METALLOTHIONEIN (MT).//0.99:14:64//PERCA FLUVIATILIS (PERCH).//P52725

F-HEMBB1000883//HYPOTHETICAL 7.8 KD PROTEIN (ORF62).//0.34:60:33//GUILLARDIA THETA (CRYPTO-MONAS PHI).//O78459

F-HEMBB1000887//HISTIDINE-RICH, METAL BINDING POLYPEPTIDE.//1.0:26:42//HELICOBACTER PYLORI (CAMPYLOBACTER PYLORI).//Q48251

F-HEMBB1000888

F-HEMBB1000890

F-HEMBB1000893

F-HEMBB1000908//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.0074:45:51//HOMO SAPIENS (HUMAN).// P39188

F-HEMBB1000910//PROBABLE E5 PROTEIN.//1.0.49:36//HUMAN PAPILLOMAVIRUS TYPE 58.//P26552 F-HEMBB1000913//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.29:56:46//HOMO SAPIENS (HUMAN).// P39195

F-HEMBB1000915//CYTOCHROME B (EC 1.10.2.2).//2.5e-24.62:90//HOMO SAPIENS (HUMAN).//P00156
F-HEMBB1000917//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/5.9e-26:53:66//HOMO SAPIENS (HUMAN).//
P30193

F-HEMBB1000927//NEURONAL CALCIUM SENSOR 1 (NCS-1) (FREQUENIN).//3.9e-44:182:45//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//Q91614

F-HEMBB1000947//SMALL PROLINE-RICH PROTEIN 2-1-//0.24:69:27//HOMO SAPIENS (HUMAN).//P35326
5-HEMBB1000959//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/3.0e-31:89:68//HOMO SAPIENS (HUMAN).//
P39195

F-HEMBB1000973//CONNECTIVE TISSUE GROWTH FACTOR PRECURSOR.//0.96:66:36//BOS TAURUS (BO-VINE).//018739

- F-HEMBB1000975//HISTIDINE-RICH GLYCOPROTEIN PRECURSOR (HISTIDINE-PROLINE RICH GLYCO-PROTEIN) (HPRG).//0.00042:77:41//HOMO SAPIENS (HUMAN).//P04196
- F-HEMBB1000981
- F-HEMBB1000985//MIPP PROTEIN (MURINE IAP-PROMOTED PLACENTA-EXPRESSED PROTEIN).//1.0e-18: 178:30//MUS MUSCULUS (MOUSE).//P28575

  - F-HEMBB1000996//HYPOTHETICAL 10.1 KD PROTEIN IN RHSD-GCL INTERGENIC REGION (ORFD3).//0.58: 34:35//ESCHERICHIA COLL//P33669
- F-HEMBB1001004//PROBABLE E4 PROTEIN.//0.24:110:35//HUMAN PAPILLOMAVIRUS TYPE 5B.//P26550
- 10 F-HEMBB1001008 F-HEMBB1001011//ZINC FINGER PROTEIN 7 (ZINC FINGER PROTEIN KOX4) (ZINC FINGER PROTEIN HF. 16).//3.2e-17:104:47//HOMO SAPIENS (HUMAN).//P17097
  - F-HEMBB1001014//FOTAXIN PRECURSOR (FOSINOPHIL CHEMOTACTIC PROTEIN).//1.0:58:39//RATTUS NORVEGICUS (RAT) //P97545
- F-HEMBB1001020//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.4e-07:36:75//HOMO SAPIENS (HUMAN).// 15 P39189
  - F-HEMBB1001024
    - F-HEMBB1001037//FERREDOXIN.//1.0:52:25//MOORELLA THERMOACETICA (CLOSTRIDIUM THER-MOACETICUM).//P00203
- 20 F-HEMBB1001047
  - F-HEMBB1001051//PROTEIN FAN (FACTOR ASSOCIATED WITH N-SMASE ACTIVATION).//3.4e-21:50:100// HOMO SAPIENS (HUMAN),//Q92636
  - F-HEMBB1001056//HYPOTHETICAL 29.3 KD PROTEIN (ORF92),//0.0099:115:35//ORGYIA PSEUDOTSUGATA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//O10341
- 25 F-HEMBB1001058//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!/1.1e-33:95:76//HOMO SAPIENS (HUMAN).// P39192
  - F-HEMBB1001060//HYPOTHETICAL 8.2 KD PROTEIN ZC21.7 IN CHROMOSOME III.//1.0:38:36// CAENORHABDITIS ELEGANS.//P34591 F-HEMBB1001063
- 30 F-HEMBB1001068

- - F-HEMBB1001096//NOXIUSTOXIN (NTX) (TOXIN II.11).//0.99:36:38//CENTRUROIDES NOXIUS (MEXICAN SCORPION) //P08815
  - F-HEMBB1001102//HYPOTHETICAL 72.5 KD PROTEIN C2F7.10 IN CHROMOSOME I.//1.1e-27:115:36// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST),//Q09701
- 35 F-HEMBB1001105//CLASS II HISTOCOMPATIBILITY ANTIGEN, M ALPHA CHAIN PRECURSOR, //0.80:70:40// HOMO SAPIENS (HUMAN),//P28067
  - F-HEMBB1001112//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//1.1e-126:287:85//RATTUS NORVEGICUS (RAT) //P38378
- F-HEMBB1001114//HYPOTHETICAL 9.6 KD PROTEIN (ORF2),//0.84:62:27//BACTERIOPHAGE L2.//P42537 40 F-HEMBB1001117
  - F-HEMBB1001119//COLLAGEN ALPHA 1(XII) CHAIN PRECURSOR.//1.6e-21:50:98//HOMO SAPIENS (HU-
    - F-HEMBB1001126//HYPOTHETICAL 55.9 KD PROTEIN EEED8.6 IN CHROMOSOME II.//1.7e-50:184:53// CAENORHARDITIS ELEGANS //009296
- F-HEMBB1001133//!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.4e-09:53:62//HOMO SAPIENS (HUMAN).// 45
  - F-HEMBB1001137//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RPB1) (FRAG-MENT),//2.0e-05:206:27//CRICETULUS GRISEUS (CHINESE HAMSTER),//P11414
    - F-HEMBB1001142///III ALU SUBFAMILY SP WARNING ENTRY IIII//4.1e-05:46:56//HOMO SAPIENS (HUMAN).// P39193
    - F-HEMBB1001151//HYPOTHETICAL 33.5 KD PROTEIN C1D4.02C IN CHROMOSOME I.//2.3e-23:109:44// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST), //Q10149
- F-HEMBB1001153//PROCOLLAGEN ALPHA 2(IV) CHAIN PRECURSOR.//0.75:76:34//ASCARIS SUUM (PIG ROUNDWORM) (ASCARIS LUMBRICOIDES),//P27393
- 55 F-HEMBB1001169//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.4e-16:71:59//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBB1001175//ANKYRIN.//3.2e-12:169:31//MUS MUSCULUS (MOUSE).//Q02357
  - F-HEMBB1001177//PERIODIC TRYPTOPHAN PROTEIN 2 HOMOLOG.//9.4e-07:148:27//HOMO SAPIENS (HU-

MAN).//Q15269

F-HEMBB1001182//HYPOTHETICAL 36.0 KD PROTEIN.//1.3e-09:110:31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P54858

F-HEMBB1001199

F-HEMBB1001208//HYPOTHETICAL PROTEIN LAMBDA-SP5.//0.053:23:47//MUS MUSCULUS (MOUSE).// P15974

F-HEMBB1001209

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F-HEMBB1001210//HYPOTHETICAL PROTEIN LAMBDA-SP5.//0.14:40:37//MUS MUSCULUS (MOUSE).// P15974

10 F-HEMBB1001218/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.4e-19:49:67//HOMO SAPIENS (HU-MAN).//P39194

F-HEMBB1001221//CYTOCHROME C OXIDASE POLYPEPTIDE VIIA-LIVER PRECURSOR (EC 1.9.3.1).//0.11: 44:38/HOMO SAPIENS (HUMAN).//P14406

F-HEMBB1001234//65 KD YES-ASSOCIATED PROTEIN (YAP65).//2.0e-45:192:53//MUS MUSCULUS (MOUSE).//P46938

F-HEMBB1001242/HYPOTHETICAL 143.3 KD TRP-ASP REPEATS CONTAINING PROTEIN C12G12.13C IN CHROMOSOME I./I5.5e-37:226:41//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09876

F-HEMBB1001249//OXALOACETATE DECARBOXYLASE GAMMA CHAIN (EC 4.1.1.3).//1.0:23:43//KLEBSIEL-LA PNEUMONIAE.//P13155

F-HEMBB1001253//METALLOTHIONEIN-IH (MT-1H) (METALLOTHIONEIN-0) (MT-0).//0.14:16:43//HOMO SA-PIENS (HUMAN).//P80294

F-HEMBB1001254///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/1.4e-12:40:75//HOMO SAPIENS (HUMAN).// P39195

F-HEMBB1001267//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.0e-12:33:78//HOMO SAPIENS (HUMAN).//

F-HEMBB1001271//HYPOTHETICAL 25.1 KD PROTEIN B0302.5 IN CHROMOSOME X.//1.0:58:37// CAENORHABDITIS ELEGANS.//Q10928

F-HEMBB1001282//ANKYRIN HOMOLOG PRECURSOR.//9.5e-13:206:31//CHROMATIUM VINOSUM.//Q06527 F-HEMBB1001288//COPPER HOMEOSTASIS PROTEIN CUTC.//4.6e-42:163:51//ESCHERICHIA COLI.//

F-HEMBB1001289//HYPOTHETICAL PROTEIN ORF-1137.//1.0e-05:106:26//MUS MUSCULUS (MOUSE).// P11260

F-HEMBB1001294//GTP-BINDING PROTEIN TC10.//1.3e-34:58:94//HOMO SAPIENS (HUMAN).//P17081

F-HEMBB1001302//HOMEOBOX PROTEIN CDX-2 (CAUDAL-TYPE HOMEOBOX PROTEIN 2) (CDX-3).//0.24: 49:46//HOMO SAPIENS (HUMAN).//Q99626

F-HEMBB1001304//GLYCINE-RICH CELL WALL STRUCTURAL PROTEIN (CLONE W10-1) (FRAGMENT).//1.0: 17:70//LYCOPERSICON ESCULENTUM (TOMATO).//001157

F-HEMBB1001314//SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.21:104:27//DROSOPHILA ERECTA (FRUIT FLY).//P13730

40 F-HEMBB1001315//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.3e-24:53:71//HOMO SAPIENS (HUMAN).// P39195

F-HEMBB1001317//HYPOTHETICAL 85.7 KD PROTEIN C13G6.03 IN CHROMOSOME 1.//0.24:90:31// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09782

F-HEMBB1001326/HYPOTHETICAL PROTEIN LAMBDA-SP5.//0.36-26:50//MUS MUSCULUS (MOUSE).//
45 P15974
F-HEMBB1001331/HYPOTHETICAL BHLF1 PROTEIN.//1.0-127:33//EPSTEIN-BARR VIRUS (STRAIN B95-8)

(HUMAN HERPESVIRUS 4).//P03181

F-HEMBB1001335//ESCARGOT/SNAIL PROTEIN HOMOLOG (FRAGMENT).//0.85;44:29//SCIARA CO-

PROPHILA (FUNGUS GNAT).//Q01799

F-HEMBB1001337//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!/4.2e-20:62:62/HOMO SAPIENS (HU-MAN)//P39194

F-HEMBB10013339/HYPOTHETICAL 17.3 KD PROTEIN CY1A11.16C.//8.2e-07:123:34//MYCOBACTERIUM

TUBERCULOSIS://Q50606

F-HEMBB1001346//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.2e-14:60:45//HOMO SAPIENS (HU-55 MAN).//P08547

F-HEMBB1001348//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.6e-14:61:62//HOMO SAPIENS (HUMAN).// P39188

F-HEMBB1001356

F-HEMBB1001364

F-HEMBB1001366/HISTIDINE-RICH PROTEIN.//0.87:26:42//PLASMODIUM FALCIPARUM (ISOLATE FCM17 / SENEGAL).//P14586

F-HEMBB1001367//IIII ALU SUBFAMILY SC WARNING ENTRY IIII//8.6e-40:146:61//HOMO SAPIENS (HU-MAN) //P39192

F-HEMBR1001369

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F-HEMBB1001380//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/1.9e-25:49:83//HOMO SAPIENS (HUMAN).//

F-HEMBB1001384/BH3 INTERACTING DOMAIN DEATH AGONIST (BID).//0.80:95:29//MUS MUSCULUS (MOUSE).///P70444

F-HEMBB1001387//PEA2 PROTEIN (PPF2 PROTEIN).//0.022:117:34//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P40091

F-HEMBB1001394//ALPHA-ADAPTIN A (CLATHRIN ASSEMBLY PROTEIN COMPLEX 2 ALPHA-A LARGE CHAIN) (100 KD COATED VESICLE PROTEIN A) (PLASMA MEMBRANE ADAPTOR HA2/AP2 ADAPTIN ALPHA

15 A SUBUNIT).//0.38:85:31//MUS MUSCULUS (MOUSE).//P17426 F-HEMBB1001410

> F-HEMBB1001424//PHOTOSYSTEM II 4 KD REACTION CENTRE PROTEIN PRECURSOR.//0.99:37:21// ORYZA SATIVA (RICE).//P12162

F-HEMBB1001426//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.0035:40:60//HOMO SAPIENS (HUMAN).// P39195

F-HEMBB1001429/(CYTOSOL AMINOPEPTIDASE) (EC 3.4.11.1) (LEUCINE AMINOPEPTIDASE) (LAP) (LEU-CYL AMINOPEPTIDASE) (PROLINE AMINOPEPTIDASE) (EC 3.4.11.5) (PROLYL AMINOPEPTIDASE).//1.1e-92/1:36//BOS TAURUS (BOVINE).//P00727

F-HEMBB1001436//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//3.4e-30:57:78//HOMO SAPIENS (HUMAN).//

F-HEMBB1001443/JPYRUVATE DEHYDROGENASE (LIPOAMIDE);PHOSPHATASE PRECURSOR (PDP) (EC 3.1.3.43) (PVUVATE DEHYDROGENASE PHOSPHATASE, CATALYTIC SUBUNIT (PDPC) //2.5e-79.155-97// BOS TAURUS (BOVINE).//P35816

E-HFMBB10014449

90 F-HEMBB1001454//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE M) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//1.1e-05:196:31//HOMO SAPIENS (HUMAN).//P10161

F-HEMBB1001458//24 KD ANTIGEN (FRAGMENT).//0.94:18:50//PLASMODIUM CHABAUDI.//P14592

F-HEMBB1001464//PPF2L ANTIGEN (FRAGMENT).//1.0:45:28//PLASMODIUM FALCIPARUM (ISOLATE PALO ALTO / UGANDA).//P07765

F-HEMBB1001482//GASTRULA ZINC FINGER PROTEIN XLCGF16.1 (FRAGMENT).//4.2e-10:37:43//XENO-PUS LAEVIS (AFRICAN CLAWED FROG).//P18712 F-HEMBB1001500

F-HEMBB1001521//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.4e-39:59:72//HOMO SAPIENS (HUMAN).// P39188

40 F-HEMBB1001527//HOMEOBOX PROTEIN HOX-B5 (XLHBOX-4) (XHOX-1B) (FRAGMENT).//0.21:131:25// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P09019

F-HEMBB1001531//GENE 32 PROTEIN (GP32).//0.88:95:30//MYCOBACTERIOPHAGE L5.//Q05241

F-HEMBB1001535//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:31:38//LUMBRICUS TERRESTRIS (COMMON EARTHWORM).//Q34942

45 F-HEMBB1001536

F-HEMBB1001537//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//0.0063:52:50//HOMO SAPIENS (HU-MAN) //P39191

F-HEMBB1001555//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.9e-23:69:63//HOMO SAPIENS (HUMAN).//

F-HEMBB1001562//RABPHILIN-3A //0 087:147:27//RATTUS NORVEGICUS (RAT) //P47709

F-HEMBB1001564//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//5.9e-27:107:54//HOMO SAPIENS (HUMAN).//P08547

F-HEMBB1001565//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.9e-12:51:54//HOMO SAPIENS (HU-MAN).//P39194

55 F-HEMBB1001585

F-HEMBB1001586

F-HEMBB1001588//HYPOTHETICAL 12.3 KD PROTEIN IN GAP1-NAP1 INTERGENIC REGION.//0.0031:31:48// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36140

- F-HEMBB1001603
- F-HEMBB1001618//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASEL//0.00076;47:44//MUS MUSCULUS (MOUSE).//P11369
- F-HEMBB1001619//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//1.0:52:32//HOMO SAPIENS (HUMAN).//P22531
  - F-HEMBB1001630
    - F-HEMBB1001635//METALLOTHIONEIN-LIKE PROTEIN TYPE 2 A.//1.0:27:44//LYCOPERSICON ESCULEN-TUM (TOMATO) //Q40157
- F-HEMBB1001637//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.0042:26:73//HOMO SAPIENS (HUMAN).//
- 10 P39188 F-HEMBB1001641

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- F-HEMBB1001653//SURVIVAL MOTOR NEURON PROTEIN 1.//0.51:36:47//CANIS FAMILIARIS (DOG).//
- F-HEMBB1001665//HOMEOBOX PROTEIN ENGRAILED-1 (HU-EN-1).//0.0030:135:34//HOMO SAPIENS (HU-MAN).//Q05925
- MAN).//Q05925 F-HEMBB1001668//PROBABLE 60S RIBOSOMAL PROTEIN L39.//0.99:25:44//CAENORHABDITIS ELEGANS.//
  - P52814
    F-HEMBB1001673//HYPOTHETICAL 46.1 KD PROTEIN IN ERP5-ORC6 INTERGENIC REGION.//0.0054:128:
  - 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38823. F-HEMBB1001684//SUPPRESSOR PROTEIN SRP40.//0.56:81:34//SACCHAROMYCES CEREVISIAE (BAK-
- ER'S YEAST).//P32583
  - F-HEMBB1001685//CYTOCHROME C OXIDASE POLYPEPTIDE VIII-HEART PRECURSOR (EC 1.9.3.1) (VIIIB) (IX).//1.0:21:47//BOS TAURUS (BOVINE).//P10175
- F-HEMBB1001695//MYOSIN IC HEAVY CHAIN.//8.9e-05:86:40//ACANTHAMOEBA CASTELLANII (AMOEBA).//
  25 P10569
  - F-HEMBB1001704//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//9.0e-08:35:71//HOMO SAPIENS (HUMAN).// P39195
  - F-HEMBB1001706//CORNIFIN (SMALL PROLINE-RICH PROTEIN I) (SPR-I) (SMALL PROLINE-RICH SQUA-MOUS CELL MARKER) (SPRP).//0.91:39:41//SUS SCROFA (PIG).//P35323
- 30 F-HEMBB1001707//FERREDOXIN-LIKE PROTEIN IN NIF REGION.//1.0:43:23//BRADYRHIZOBIUM JAPONI-CUM.//P27394
  - F-HEMBB1001717//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4 (EC 1.6.5.3) (FRAGMENT).//1.0:71:25//
    LEMUR CATTA (RING-TAILED LEMUR).//Q34878
    - F-HEMBB1001735///// ALU SUBFAMILY SQ WARNING ENTRY !!!!//9.0e-35:97:74//HOMO SAPIENS (HU-MAN) //P39194
      - F-HEMBB1001736//EUKARYOTIC TRANSLATION INITIATION FACTOR 3 BETA SUBUNIT (EIF-3 BETA) (EIF3 P116) (EIF3 P110).//0.00069:180:28//HOMO SAPIENS (HUMAN).//P55884
    - F-HEMBB1001747
- F-HEMBB1001749///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.8e-43:75:70//HOMO SAPIENS (HUMAN).//
  40 P39195
- F-HEMBB1001753//PROTEIN Q300.//0.00091:16:81//MUS MUSCULUS (MOUSE).//Q02722
  - F-HEMBB1001756//CYCLIN-DEPENDENT KINASES REGULATORY SUBUNIT 2 (XE-P9).//0.94:35:42//XENO-PUS LAEVIS (AFRICAN CLAWED FROG).//(Q91879 F-HFMBB1001760
- 45 F-HEMBB1001762//GENE 35 PROTEIN (GP35).//0.76:21:47//MYCOBACTERIOPHAGE L5.//Q05245
- F-HEMBB1001785
  - F-HEMBB1001797//CHLOROPLAST 50S RIBOSOMAL PROTEIN L35.//0.99:41:31//PORPHYRA PURPUREA.// P51270
  - F-HEMBB1001802
- F-HEMBB1001812//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.2e-39:54:77//HOMO SAPIENS (HUMAN).// P39193
  - F-HEMBB1001816//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//1.1e-19:97:57//HOMO SAPIENS (HU-MAN).//P39194
- F-HEMBB1001831//HYPOTHETICAL 45.6 KD PROTEIN IN COX5A-ALG11 INTERGENIC REGION.//0.62:204: 23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53951
- F-HEMBB1001834//GLYCINE-RICH RNA-BINDING PROTEIN 1 (FRAGMENT).//0.0014:40:45//SORGHUM VUL-GARE (SORGHUM).//Q99069
  - F-HEMBB1001836//IIII ALU SUBFAMILY SB2 WARNING ENTRY !!!!//7.1e-14:85:61//HOMO SAPIENS (HU-

MAN).//P39191

F-HEMBB1001839//PROBABLE E4 PROTEIN.//0.61:49:34//HUMAN PAPILLOMAVIRUS TYPE 6C.//P20969 F-HEMBB1001850

F-HEMBB1001863//!!! ALU SUBFAMILY SO WARNING ENTRY !!!!//2.7e-30:57:68//HOMO SAPIENS (HU-

MAN) //P39194 F-HEMBB1001867

F-HEMBB1001868//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).// 0.00036:47:53//NICOTIANA TABACUM (COMMON TOBACCO).//P13983

F-HEMBB1001869//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.0e-11:95:45//HOMO SAPIENS (HUMAN).//

P30188 F-HEMBB1001872//HYPOTHETICAL 8.2 KD PROTEIN IN LEE8-EP INTERGENIC REGION //1.0:34:38//

AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41459 F-HEMBB1001874

F-HEMBB1001875

15 F-HEMBR1001880

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F-HEMBB1001899//GENE 11 PROTEIN.//1.0:45:31//SPIROPLASMA VIRUS SPV1-R8A2 B.//P15902

F-HEMBB1001905//HYPOTHETICAL 81.7 KD PROTEIN IN MOL1-NAT2 INTERGENIC REGION.//8.8e-54:216: 51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P48234

F-HEMBB1001906

20 F-HEMBB1001908//MONOCYTIC LEUKEMIA ZINC FINGER PROTEIN.//6.3e-51:138:80//HOMO SAPIENS (HU-MAN).//Q92794

F-HEMBB1001910

F-HEMBB1001911

F-HEMBB1001915//UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 64E (EC 3.1.2.15) (UBIQUITIN THI-25 OLESTERASE 64E) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 64E) (DEUBIQUITINATING ENZYME 64E).//2.3e-27:71:70//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q24574

F-HEMBB1001921//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//9.8e-13:75:53//HOMO SAPIENS (HU-MAN).//P08547

F-HEMBB1001922

F-HEMBB1001925//EPITHELIAL MEMBRANE PROTEIN-1 (EMP-1) (TUMOR-ASSOCIATED MEMBRANE PRO-TEIN).//1.0:55:30//MUS MUSCULUS (MOUSE).//P47801

F-HEMBB1001930//HYPOTHETICAL 9.6 KD PROTEIN K10D2.7 IN CHROMOSOME III.//0.43:49:26// CAENORHARDITIS ELEGANS //009412

F-HEMBB1001944///III ALU SUBFAMILY SB WARNING ENTRY !!!!//5.1e-34:63:85//HOMO SAPIENS (HUMAN).//

F-HEMBB1001945//NONSPECIFIC LIPID-TRANSFER PROTEIN (LTP) (PHOSPHOLIPID TRANSFER PRO-TEIN) (PLTP).//0.28:45:40//AMARANTHUS CAUDATUS (LOVE-LIES-BLEEDING) (INCA-WHEAT).//P80450

F-HEMBB1001947//PROTEIN UL24.//0.48:42:47//HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN 17).//P10208 F-HEMBB1001950//HYPOTHETICAL 42.6 KD PROTEIN IN GSHB-ANSB INTERGENIC REGION (O378).//1.6e-

F-HFMBB1001952

F-HEMBB1001953

24:162:36//ESCHERICHIA COLI.//P52062

F-HEMBB1001957//!!! ALU SUBFAMILY J WARNING ENTRY !!!!/2.7e-11:51:60//HOMO SAPIENS (HUMAN).//

45 F-HEMBB1001962//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.6e-24:163:42//HOMO SAPIENS (HUMAN).// P30188

F-HEMBB1001967//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.1e-35:55:80//HOMO SAPIENS (HUMAN).//

F-HEMBB1001973///III ALU SUBFAMILY SC WARNING ENTRY IIII//2.1e-37:108:75//HOMO SAPIENS (HU-MAN) //P39192

F-HEMBB1001983//LYSIS PROTEIN (E PROTEIN) (GPE).//0.84:45:37//BACTERIOPHAGE ALPHA-3.//P31280 F-HEMBR1001988

F-HEMBB1001990

F-HEMBB1001996//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.4e-14:98:40//HOMO SAPIENS (HU-

F-HEMBB1001997//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.1e-19:38:73//HOMO SAPIENS (HUMAN).//

F-HEMBB1002002//CYTOCHROME C BIOGENESIS PROTEIN CCSA //1.0:150:25//PORPHYRA PURPUREA.//

### P51369

F-HEMBB1002005//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//7.6e-12:94:40//HOMO SAPIENS (HUMAN).//

# F-HEMBB1002009

- F-HEMBB1002015//HYPOTHETICAL 7.7 KD PROTEIN IN MRR-TSR INTERGENIC REGION (F67).//1.0:17:47// ESCHERICHIA COLL.//P39395
  - F-HEMBB1002042//CYTOCHROME P450 4C1 (EC 1.14.14.1) (CYPIVC1).//2.4e-50:139:55//BLABERUS DISCOIDALIS (TROPICAL COCKROACH).//P29981
  - F-HEMBB1002043//HYPOTHETICAL 9.5 KD PROTEIN IN DHFR 3'REGION (ORF3).//0.052:40:42//HERPESVI-
- RUS SAIMIRI (SUBGROUP C / STRAIN 488)./P22577
  F-HEMBB1002044//CELLULOSE COMPLEMENTING PROTEIN.//0.45:87:33//ACETOBACTER XYLINUM (ACETOBACTER PASTEURIANUS)./P37697
  - F-HEMBB1002045//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.78:18:55//HOMO SAPIENS (HUMAN).//

### 15 F-HEMBR1002049

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- F-HEMBB1002050//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONE CP7) [CONTAINS: BASIC PEPTIDE P-FI (FRAGMENT).//1.09-06:188:27//HOMO SAPIENS (HUMAN).//P02812
  - F-HEMBB1002068//HOMEOBOX PROTEIN HOX-A4 (CHOX-1.4).//0.0023:56:44//GALLUS GALLUS (CHICK-FN)//P17277
- 20 F-HEMBB1002069//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).// 0.0074:134:33//NICOTIANA TABACUM (COMMON TOBACCO).//P13983
  - F-HEMB91020992/IENV POLYPROTEIN PRECURSOR (COAT POLYPROTEIN) [CONTAINS: OUTER MEM-BRANE PROTEIN GP70; TRANSMEMBRANE PROTEIN P20E].//2.4e-07:75-40//BABOON ENDOGENOUS VI-RUS (STRAIN M7).//P10269
- 25 F-HEMBB1002094/IIII ALU SUBFAMILY SB2 WARNING ENTRY IIII//1.9e-24:63:82//HOMO SAPIENS (HU-MAN).//P39191
  - F-HEMBB1002115//EC PROTEIN HOMOLOG (ZINC-METALLOTHIONEIN CLASS II).//0.94:26:42//ZEA MAYS (MAIZE).//P43401
- F-HEMBB1002134//ZINC-FINGER PROTEIN NEURO-D4.//4.6e-57:176:67//RATTUS NORVEGICUS (RAT).//
  P56163
- F-HEMBB1002139//CHLOROPLAST 50S RIBOSOMAL PROTEIN L35.//1.0:17:52//PORPHYRA PURPUREA.// P51270
  - F-HEMBB1002142//EARLY NODULIN 20 PRECURSOR (N-20).//0.087:52:36//MEDICAGO TRUNCATULA (BAR-REL MEDIC).//P93329
- 35 F-HEMBB1002152//HYPOTHETICAL 12.3 KD PROTEIN IN RPL3-RPL33 INTERGENIC REGION (ORF102).// 5.8e-05:61:37//CYANOPHORA PARADOXA.//P15811
  - F-HEMBB1002189//HYPOTHETICAL PROTEIN UL125.//1.0:77:32//HUMAN CYTOMEGALOVIRUS (STRAIN AD169\)//P16835

### F-HEMBB1002190

- F-HEMBB1002193/ITYROSINE-PROTEIN KINASE RECEPTOR TYRO3 PRECURSOR (TYROSINE-PROTEIN KINASE RSE) (TYROSINE-PROTEIN KINASE SKY) (TYROSINE-PROTEIN KINASE DTK).//1.2e-27:59:100// HOMO SAPIENS (HUMAN)./006418
  - F-HEMBB1002217//ZINC FINGER PROTEIN 184 (FRAGMENT).//6.6e-22:106:50//HOMO SAPIENS (HUMAN).//
- 45 F-HEMBB1002218//PROTEIN Q300.//0.85:19:52//MUS MUSCULUS (MOUSE).//Q02722
  - F-HEMBB1002232//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//9.6e-21:56:71//HOMO SAPIENS (HUMAN).// P39195

### F-HEMBB1002247

- F-HEMBB1002249/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.2e-29:93:69//HOMO SAPIENS (HU-MAN) //P39194
- F-HEMBB002254//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.0e-29:101:67//HOMO SAPIENS (HU-MAN).//P39194
  - F-HEMBB1002255//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 3 (EC 1.6.5.3),//1.0:73:28//PARA-MECIUM TETRAUREI IA //P15579
- 55 F-HEMBB1002266//GLUTAMIC ACID-RICH PROTEIN PRECURSOR.//0.0079:151:26//PLASMODIUM FALCI-PARUM (ISOLATE FC27 / PAPUA NEW GUINEA).//P13816
  - F-HEMBB1002280//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//5.2e-15:182:36//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548

F-HEMBB1002300

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F-HEMBB1002306//!!!! ALU SUBFAMILY SX WARNING ENTRY!!!!//0.00011:26:84//HOMO SAPIENS (HU-MAN).//P39195

F-HEMBB1002327//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//4.1e-11:41:85//HOMO SAPIENS (HUMAN).// P39189

F-HEMBB1002329//HYPOTHETICAL 74.0 KD PROTEIN IN CAJ1-HOM3 INTERGENIC REGION.//9.9e-17:232: 30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40032

F-HEMBB1002340
F-HEMBB1002342//HYPOTHETICAL 32.5 KD PROTEIN IN MSH6-BMH2 INTERGENIC REGION.//3.6e-40:102:

57//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q03835
F-HEMBE1002358//THYMIDYLATE KINASE (EC 2.7.4.9) (DTMP KINASE).//6.1e-30:63:96//HOMO SAPIENS (HUMAN).//P23919

(HUMAN),/P23979
F-HEMBB1002359/HYPOTHETICAL 7.1 KD PROTEIN C6G9.01C IN CHROMOSOME I.//0.97:28:46//
SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//092346

15 F-HEMBB1002364//RETROVIRUS-RELATED POL POLYPROTEIN (FRAGMENT).//0.47:119:25//HOMO SAPI-ENS (HUMAN).//P12895

F-HEMBB1002371//HYPOTHETICAL 15.5 KD PROTEIN C2F7.12 IN CHROMOSOME I PRECURSOR.//3.0e-05: 111:30//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09703

F-HEMBB1002381//PUTATIVE CUTICLE COLLAGEN C09G5.4.//0.34:105:34//CAENORHABDITIS ELEGANS.//

F-HEMBB1002383//ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6).//0.049:103:32//AQUIFEX AEOLI-CUS.//086586

F-HEMBB1002387//10 KD CHAPERONIN (PROTEIN CPN10) (PROTEIN GROES) (HEAT SHOCK PROTEIN 11).//0.18:75:28//RICKETTSIA TSUTSUGAMUSHI.//P16626

25 F-HEMBB1002409//HIGH MOBILITY GROUP PROTEIN HMG-Y.//0.014:81:36//MUS MUSCULUS (MOUSE).// P17095

F-HEMBB1002415

F-HEMBB1002425/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.8e-18:55:70//HOMO SAPIENS (HU-MAN).//P39194

30 F-HEMBB1002442//LIN-10 PROTEIN.//5.1e-15:121:31//CAENORHABDITIS ELEGANS.//P34692

F-HEMBB1002453//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.2e-32:54:75//HOMO SAPIENS (HUMAN).// P39189

F-HEMBB1002457//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.5e-07:31:64//HOMO SAPIENS (HUMAN).// P39188

35 F-HEMBB1002458//MALE SPECIFIC SPERM PROTEIN MST84DA.//0.92:28:53//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01642

F-HEMBB1002477//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RPB1) (FRAGMENT).//0.0066:198:27//CRICETULUS GRISEUS (CHINESE HAMSTER).//P11414

F-HEMBB1002489//SPLICEOSOME ASSOCIATED PROTEIN 49 (SAP 49) (SF3B53).//0.030:182:28//HOMO SA-40 PIENS (HUMAN).//Q15427

F-HEMBB1002492

F-HEMBB1002495//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//2.1e-08:41:75//HOMO SAPIENS (HUMAN).// P39192

F-HEMBB1002502//RETROVIRUS-RELATED POL POLYPROTEIN (FRAGMENT).//0.00030:31:77//HOMO SA-45 PIENS (HUMAN).//P12895

F-HEMBB1002509

F-HEMBB1002510

F-HEMBB1002520//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.8e-36:162:50//NYCTICEBUS COUCANG (SLOW LORIS).//P08548

F-HEMBB1002522//7 KD PROTEIN (ORF 4).//0.77:32:40//CHRYSANTHEMUM VIRUS B (CVB).//P37990

F-HEMBB1002534///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//3.1e-36:80:73//HOMO SAPIENS (HUMAN).// P39195

E-HEMBR1002545

55 F-HEMBB1002550//HOMEOBOX PROTEIN HOX-D11 (HOX-4.6) (HOX-5.5).//3.8e-05:83:34//MUS MUSCULUS (MOUSE).//P23813

F-HEMBB1002556

F-HEMBB1002579//SPLICING FACTOR U2AF 35 KD SUBUNIT (U2 AUXILIARY FACTOR 35 KD SUBUNIT) (U2

- SNRNP AUXILIARY FACTOR SMALL SUBUNIT) (FRAGMENT) //5.0e-06:27:77//SUS SCROFA (PIG) //029350 F-HEMBB1002582//PROTEINASE INHIBITOR://1.0:27:40//SOLANUM MELONGENA (EGGPLANT) (AUBER-GINE).//P01078
- F-HEMBB1002590//HYPOTHETICAL PROTEIN IN MMSB 3'REGION (ORF1) (FRAGMENT).//1.9e-20:90:54// PSELIDOMONAS AERLIGINOSA //P28812

  - F-HEMBB1002600//NOVEL ANTIGEN 2 (NAG-2).//1.9e-60:187:59//HOMO SAPIENS (HUMAN).//O14817
  - F-HEMBB1002601//M PROTEIN, SEROTYPE 6 PRECURSOR.//1.0:71:35//STREPTOCOCCUS PYOGENES.// PARARA
- 10 F-HEMBB1002603

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- F-HEMBB1002607//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT),//0.0032:142:33//HOMO SAPIENS (HUMAN),//P10162
- F-HEMBB1002610//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG //2.0e-11:79:49//HOMO SAPIENS (HU-MAN).//P08547
- 15 F-HEMBB1002613//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.9e-08:41:60//HOMO SAPIENS (HUMAN).//
  - F-HEMBB1002614//HYPOTHETICAL 9.5 KD PROTEIN.//1.0:40:35//VACCINIA VIRUS (STRAIN COPENHA-GEN) //P20553
  - F-HEMBB1002617/INSECT TOXIN 1 (BOT IT1).//1.0:44:29//BUTHUS OCCITANUS TUNETANUS (COMMON EUROPEAN SCORPION) //P55902
  - F-HEMBB1002623//HYPOTHETICAL 9.7 KD PROTEIN (ORF88) (PUTATIVE DNA-BINDING PROTEIN),//0.42: 31:54//BACTERIOPHAGE P4.//P12552
    - F-HEMBB1002635//STRESS-ACTIVATED PROTEIN KINASE JNK3 (EC 2.7.1.-) (C-JUN N-TERMINAL KINASE (MAP KINASE P49 3F12).//6.2e-17:44:95//HOMO SAPIENS (HUMAN).//P53779
- 25 F-HEMBB1002664//SMALL NUCLEAR RIBONUCLEOPROTEIN ASSOCIATED PROTEIN B (SM-B) (SNRNP-B) (SM11) (FRAGMENT), //1.0:57:36//RATTUS NORVEGICUS (RAT), //P17136
  - F-HEMBB1002677//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.9e-06:194:34//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548
- F-HEMBB1002683//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3) (FRAGMENT).//0.96:56: 35//LEMUR CATTA (RING-TAILED LEMUR).//Q34879
  - F-HEMBB1002684//SILLUCIN.//1.0:16:50//RHIZOMUCOR PUSILLUS.//P02885
    - F-HEMBB1002686
    - F-HEMBB1002692
    - F-HEMBB1002697//HELIX-DESTABILIZING PROTEIN (SINGLE-STRANDED DNA BINDING PROTEIN) (GPV).// 0.57:36:38//BACTERIOPHAGE FD, BACTERIOPHAGE F1, AND BACTERIOPHAGE M13.//P03669
- F-HEMBR1002699
  - F-HEMBB1002702 F-HEMBB1002712
  - F-HEMBB1002705//HYPOTHETICAL 34.8 KD PROTEIN C4H3.04C IN CHROMOSOME I.//3.6e-40:180:37// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10212
- - F-MAMMA1000009//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!/1.2e-32:95:75//HOMO SAPIENS (HU-MAN).//P39189
- F-MAMMA1000019
- F-MAMMA1000020//DIMETHYLANILINE MONOOXYGENASE IN-OXIDE FORMING) 5 (EC 1.14.13.8) (HEPATIC 45 FLAVIN-CONTAINING MONOOXYGENASE 5) (FMO 5) (DIMETHYLANILINE OXIDASE 5).//5.2e-12:24:100//HO-MO SAPIENS (HUMAN) //P49326
  - F-MAMMA1000025//BETA-2-MICROGLOBULIN PRECURSOR.//1.0:73:26//BRACHYDANIO RERIO (ZE-BRAFISH) (ZEBRA DANIO),//Q04475
  - F-MAMMA1000043//HYPOTHETICAL PXBL-I PROTEIN (FRAGMENT).//0.057:130:31//BOVINE LEUKEMIA VI-RUS (JAPANESE ISOLATE BLV-1) (BLV).//P03412
  - F-MAMMA1000045
    - F-MAMMA1000055//TESTIN 2 (TES2) [CONTAINS: TESTIN 1 (TES1)].//7.5e-44:138:55//MUS MUSCULUS
- F-MAMMA1000057//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/1.2e-39:92:69//HOMO SAPIENS (HU-55 MAN).//P39194
  - F-MAMMA1000069//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//0.0044:96:34//ORGYIA PSEUDOTSUGATA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV), //O10341
    - F-MAMMA1000084//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/5.4e-28:94:73//HOMO SAPIENS (HU-

MAN).//P39195

F-MAMMA1000086/PUTATIVE CYSTEINYL-TRNA SYNTHETASE C29E6.06C (EC. 6.1.1.16) (CYSTEINE-TRNA LIGASE) (CYSRS)//6.66-38:90:51//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//009860 F-MAMMA1000092/IIII ALU SUBFAMILY SC WARNING ENTRY!!!!/6.46-30-43:86/HOMO SAPIENS (HU-

MAN).//P39192

F-MAMMA1000103//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.038:17:52//HOMO SAPIENS (HUMAN).//P22531

F-MAMMA1000117//50S RIBOSOMAL PROTEIN L24E (HL21/HL22).//0.90:25:48//HALOARCULA MARISMORTUI (HALOBACTERIUM MARISMORTUI).//P14116

10 F-MAMMA1000129//HYPOTHETICAL BHLF1 PROTEIN.//0.0016:75:40//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181

F-MAMMA1000133

F-MAMMA1000134//HYPOTHETICAL PROTEIN MJ0647.//1.0:41:41//METHANOCOCCUS JANNASCHII.// Q58063

F-MAMMA1000139//GUANINE NUCLEOTIDE-BINDING PROTEIN G(I)/G(S)/G(O) GAMMA-3 SUBUNIT.//0.99: 69:28//BOS TAURUS (BOVINE), AND MUS MUSCULUS (MOUSE).//P29798

F-MAMMA1000143//CALPAIN INHIBITOR (CALPASTATIN) (FRAGMENT).//0.023:111:27//MUS MUSCULUS (MOUSE).//P51125

F-MAMMA1000155//PUTATIVE CUTICLE COLLAGEN C09G5.5.//0.018:125:34//CAENORHABDITIS ELE-GANS.//Q09456

F-MAMMA1000163/MERCURIC TRANSPORT PROTEIN PERIPLASMIC COMPONENT PRECURSOR (PERI-PLASMIC MERCURY ION BINDING PROTEIN) (MERCURY SCAVENGER PROTEIN)./0.11:88:25//SHEWANEL-LA PUTREFACIENS (PSEUDOMONAS PUTREFACIENS)./0.54463

F-MAMMA1000171

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25 F-MAMMA1000173//DREBRIN E.//7.6e-41:197:43//HOMO SAPIENS (HUMAN).//Q16643

F-MAMMA1000175//GAMMA-THIONIN HOMOLOG PPT PRECURSOR.//0.92:39:38//PETUNIA INTEGRIFOLIA (VIOLET-FLOWERED PETUNIA) (PETUNIA INFLATA).//Q40901

F-MAMMA1000183//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2) //2.4e-106:249:61//HOMO SA-PIENS (HUMAN).//P51523

F-MAMMA1000198//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.0014:35:42//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01645

F-MAMMA1000221

F-MAMMA1000227//6.8 KD MITOCHONDRIAL PROTEOLIPID.//1.0:30:40//MUS MUSCULUS (MOUSE).// P56379

F-MAMMA1000241//PHOTOSYSTEM I REACTION CENTRE SUBUNIT X (PSI-K).//1.0:40:37//PORPHYRA PUR-PUREA.//P51370

F-MAMMA1000251//HYPOTHETICAL 6.8 KD PROTEIN IN FIC-PPIA INTERGENIC REGION.//0.99:29:48//SAL-MONELLA TYPHIMURIUM.//P37771

F-MAMMA1000254//HYPOTHETICAL 6.0 KD PROTEIN IN THI12 5'REGION.//1.0:20:50//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53820

F-MAMMA1000257//HYPOTHETICAL 50.0 KD PROTEIN IN HEML 3'REGION (ORF2).//0.22:50:44//PSEU-DOMONAS AERUGINOSA.//Q51470

F-MAMMA1000264//GASTRIN-RELEASING PEPTIDE RECEPTOR (GRP-R) (GRP-PREFERRING BOMBESIN RECEPTOR).//0.80:39:43//HOMO SAPIENS (HUMAN).//P30550

45 F-MAMMA1000266

F-MAMMA1000270//!!!! ALU SUBFAMILY SB WARNING ENTRY!!!!//9.5e-42:95:84//HOMO SAPIENS (HU-MAN).//P39189

F-MAMMA1000277//PROCOLLAGEN ALPHA 1(II) CHAIN PRECURSOR [CONTAINS: CHONDROCALCIN].//
0.0062:90:34//MUS MUSCULUS (MOUSE).//P28481

F-MAMMA1000278//C-HORDEIN (CLONE PC HOR1-3) (FRAGMENT).//0.00096:59:33//HORDEUM VULGARE (BARLEY).//P17991

F-MAMMA1000279///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//8.4e-17:56:76//HOMO SAPIENS (HU-MAN).//P39195

F-MAMMA1000284//ARYL HYDROCARBON RECEPTOR NUCLEAR TRANSLOCATOR 2 (ARNT PROTEIN 2).//
0.017:146:30//MUS MUSCULUS (MOUSE).//Q61324

F-MAMMA1000287///!!! ALU SUBFAMILY SB WARNING ENTRY!!!!//1.5e-32:84:58//HOMO SAPIENS (HU-MAN).//P39189

F-MAMMA1000302//C-HORDEIN (CLONE PC-919) (FRAGMENT).//1.0:42:33//HORDEUM VULGARE (BAR-

### LEY).//P17992

F-MAMMA1000307//PROBABLE E4 PROTEIN.//0.21:71:30//RHESUS PAPILLOMAVIRUS TYPE 1 (RHPV 1).// P24832

F-MAMMA1000309//COLLAGEN ALPHA 1(VIII) CHAIN PRECURSOR (ENDOTHELIAL COLLAGEN).//0.0026: 141:36//HOMO SAPIENS (HUMAN).//P27658

### F-MAMMA1000312

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F-MAMMA1000313//DNA REPAIR PROTEIN RADC HOMOLOG (25 KD PROTEIN) (FRAGMENT).//0.76:52:32// STAPHYLOCOCCUS AUREUS //P31337

### F-MAMMA1000331

10 F-MAMMA1000339//50S\_RIBOSOMAL\_PROTEIN\_L29P.//0.78:32:46//METHANOBACTERIUM\_THERMOAU-TOTROPHICUM //O26117

F-MAMMA1000340//HYPOTHETICAL 29.4 KD PROTEIN IN STE6-LOS1 INTERGENIC REGION.//1.0:29:58// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST). //P36039

F-MAMMA1000348//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.5e-09:63:60//HOMO SAPIENS (HUMAN).// P39188

### F-MAMMA1000356//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/3.3e-05:42:52//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1000360

F-MAMMA1000361//IIII ALU SUBFAMILY SB WARNING ENTRY IIII//44e-33:84:72//HOMO SAPIENS (HU-

F-MAMMA1000372//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/6.6e-21:53:71//HOMO SAPIENS (HU-MAN).//P39193

### F-MAMMA1000385

F-MAMMA1000388//OX40L RECEPTOR PRECURSOR (ACT35 ANTIGEN) (TAX-TRANSCRIPTIONALLY ACTI-25 VATED GLYCOPROTEIN 1 RECEPTOR) (CD134 ANTIGEN) //0.40;72;36//HOMO SAPIENS (HUMAN) //P43489 F-MAMMA1000395//RABPHILIN-3A (FRAGMENT),//0.032;125;25//MUS MUSCULUS (MOUSE),//P47708

F-MAMMA1000402//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.1e-28:266:40//HOMO SAPIENS (HU-MAN).//P08547

F-MAMMA1000410//NADH-UBIQUINONE OXIDOREDUCTASE 13 KD-B SUBUNIT (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-13KD-B) (CI-13KD-B) (B13), I/5.9e-06:32:68//HOMO SAPIENS (HUMAN), I/Q16718

F-MAMMA1000413//RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE].//6.7e-05:93:31//MUS MUSCULUS (MOUSE).//P11369

### F-MAMMA1000414

F-MAMMA1000416//HYPOTHETICAL 32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III.//4.1e-28:119:53// CAENORHARDITIS ELEGANS //009232

F-MAMMA1000421//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/2.7e-23:68:76//HOMO SAPIENS (HU-MAN).//P39194

F-MAMMA1000422//METALLOTHIONEIN (MT).//0.037:42:42//GADUS MORHUA (ATLANTIC COD).//P51902 F-MAMMA1000423

40 F-MAMMA1000424///III ALU SUBFAMILY SB WARNING ENTRY IIII/0.048:23:73//HOMO SAPIENS (HUMAN).// P39189

F-MAMMA1000429//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS5.//2.7e-05:110:30//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST),//Q92331

F-MAMMA1000431//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//9.4e-15:85:58//HOMO SAPIENS (HU-MAN).//P39194

F-MAMMA1000444//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/4.3e-25:65:76//HOMO SAPIENS (HU-MAN) //P39194

F-MAMMA1000446//ZYXIN.//0.79:155:29//GALLUS GALLUS (CHICKEN).//Q04584

F-MAMMA1000458//HYPOTHETICAL 37.7 KD PROTEIN C18B11.06 IN CHROMOSOME L//0.0048:46:43// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09713

F-MAMMA1000468//PERIOD CLOCK PROTEIN (FRAGMENT), //0.50:20:55//DROSOPHILA ROBUSTA (FRUIT FLY).//Q03296

F-MAMMA1000472//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.5e-17:106:55//HOMO SAPIENS (HUMAN),//

55 F-MAMMA1000478//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/3.9e-35:80:68//HOMO SAPIENS (HU-MAN).//P39195

F-MAMMA1000483//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.8e-24:74:77//HOMO SAPIENS (HU-MAN).//P39193

- F-MAMMA1000490//TYROSINE-PROTEIN KINASE TXK (EC 2.7.1.112) (PTK-RL-18) (RESTING LYMPHOCYTE KINASE).//0.43:21:57//MUS MUSCULUS (MOUSE).//P42682
- F-MAMMA1000500//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN).//0.61:33:54//HUMAN IM-MUNODEFICIENCY VIRUS TYPE 1 (Z2/CDC-Z34 ISOLATE) (HIV-1).//P12506
- F-MAMMA1000501///!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.1e-32:43:83//HOMO SAPIENS (HU-MAN).//P39194
  - F-MAMMA1000516
  - F-MAMMA1000522//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.0015:113:32//HOMO SAPIENS (HU-MAN).//P08547
- 1º F-MAMMA1000524//HYPOTHETICAL HOST RANGE 8.5 KD PROTEIN.//1.0:63:31//VACCINIA VIRUS (STRAIN WR).//P17359
- 15 F-MAMMA1000567//!!!! ALU SUBFAMILY SX WARNING ENTRY!!!!/5.5e-37:95:76//HOMO SAPIENS (HU-
  - F-MAMMA1000576//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//4.1e-07:34:64//HOMO SAPIENS (HU-MAN).//P39191
- F-MAMMA1000583
  F-MAMMA1000585//!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//1.5e-28:89:75//HOMO SAPIENS (HU-MAN)/P39194
  - F-MAMMA1000594///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//4.8e-24:38:71//HOMO SAPIENS (HU-MAN).//P39195
- F-MAMMA1000597//!!!! ALU SUBFAMILY SX WARNING ENTRY!!!!//1.1e-25:74:77//HOMO SAPIENS (HU-MAN).//P39195
  - F-MAMMA1000605//!!!! ALU SUBFAMILY SX WARNING ENTRY!!!!//2.1e-18:83:50//HOMO SAPIENS (HU-MAN)//P39195
  - F-MAMMA1000612//HYPOTHETICAL 34.0 KD TRP-ASP REPEATS CONTAINING PROTEIN IN SIS1-MRPL2 INTERGENIC REGION.//4.0e-42:166.48//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)///P41318
- 30 F-MAMMA1000616
  - F-MAMMA1000621

- F-MAMMA1000623//METALLOTHIONEIN-IK (MT-1K).//0.0045:25:48//HOMO SAPIENS (HUMAN).//P80296
  F-MAMMA1000629//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.00078:79:35//MUS MUSCULUS
  (MOLISE).//P05143
- 35 F-MAMMA1000643//HYPOTHETICAL 9.3 KD PROTEIN.//1.0:25:28//MAGUARI VIRUS.//P16607 F-MAMMA1000664
  - F-MAMMA1000669//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.2e-05:186:30//HOMO SAPIENS (HU-MAN).//P08547
- F-MAMMA1000670//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//1.6e-06:195:30//MUS MUSCULUS (MOUSE).//P05143
  - F-MAMMA1000672//VITELLOGENIC CARBOXYPEPTIDASE PRECURSOR (EC 3.4.16.-).//3.8e-28:184:35//
    AEDES AEGYPTI (YELLOWFEVER MOSQUITO).//P42660
  - F-MAMMA1000684//DNA-BINDING PROTEIN (VMW21).//1.1e-07:55:56//HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN 17).//P04487
- 45 F-MAMMA1000696/!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!/1.2e-31:97:74/HOMO SAPIENS (HU-MAN)./P39194 F-MAMMA1000707//METALLOTHIONEIN-II (MT-II) (MT-IIB/MT-IIA)./IO.31:19:42//CALLINECTES SAPIDUS
  - (BLUE CRAB), IP55950 F-MAMMA1000713/IXYLULOSE KINASE (EC 2.7.1.17) (XYLULOKINASE), I/1.6e-05:88:35/ILACTOBACILLUS PENTOSIUS / IP21939
  - F-ENTOSUS.III-21939

    F-MAMMA1000714I/PROTEIN-LYSINE 6-OXIDASE PRECURSOR (EC 1.4.3.13) (LYSYL OXIDASE).I/0.44:126:
    30I/FATTUS NORVEGICUS (RAT).I/P16636
    - F-MAMMA1000718//METALLOTHIONEIN-IIE (MT-2E).//1.0:51:31//ORYCTOLAGUS CUNICULUS (RABBIT).//
- 55 F-MAMMA1000720/!!!! ALU SUBFAMILY SP WARNING ENTRY!!!!//3.3e-28:60:71//HOMO SAPIENS (HU-MAN).//P39193
  - F-MAMMA1000723//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.7e-14:63:53//HOMO SAPIENS (HU-MAN).//P08547

F-MAMMA1000731//CHROMODOMAIN-HELICASE-DNA-BINDING PROTEIN 2 (CHD-2).//1.8e-43:258:43//HO-MO SAPIENS (HUMAN).//O14647

F-MAMMA1000732//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.9e-12:76:55//HOMO SAPIENS (HUMAN).// P39188

## F-MAMMA1000733

F-MAMMA1000734//NPL1 PROTEIN (SEC63 PROTEIN).//2.5e-18:181:39//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P14906

F-MAMMA1000738//HYPOTHETICAL 116.5 KD PROTEIN C20G8.09C IN CHROMOSOME I.//5.4e-52:196:58// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P87115

F-MAMMA1000744///!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!//6.3e-36:144:47//HOMO SAPIENS (HU-MAN).//P39190

F-MAMMA1000746

F-MAMMA1000752

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F-MAMMA1000760//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/6.6e-29:75:72//HOMO SAPIENS (HU-MAN) //P39195

F-MAMMA1000761//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.6e-09:59:64//HOMO SAPIENS (HU-MAN).//P39194

F-MAMMA1000775

F-MAMMA1000776//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//3.3e-35:99:74//HOMO SAPIENS (HU-MAN) //P30103

F-MAMMA1000778//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.1e-19:65:70//HOMO SAPIENS (HU-MAN)://P39195

F-MAMMA1000782

F-MAMMA1000798//HYPOTHETICAL PROTEIN ORF-1137.//0.015:59:37//MUS MUSCULUS (MOUSE).//

F-MAMMA1000802//MYOSIN IC HEAVY CHAIN.//0.35;94:41//ACANTHAMOEBA CASTELLANII (AMOEBA).//

F-MAMMA1000824//ACTIN 1.//0.046:60:31//ZEA MAYS (MAIZE).//P02582

F-MAMMA1000831//PROBABLE NI/FE-HYDROGENASÈ 1 B-TYPE CYTOCHROME SUBUNIT.//1.0:30:46//ES-CHERICHIA COLL//P19929

F-MAMMA1000839//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-28:80:58//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1000841//PUTATIVE AMIDASE (EC 3.5.1.4).//1.5e-39:130:36//METHANOBACTERIUM THERMOAU-TOTROPHICUM.//027540

35 F-MAMMA1000842//C-HORDEIN (CLONE PC-919) (FRAGMENT).//0.064:43:41//HORDEUM VULGARE (BAR-LEY).//P17992

F-MAMMA1000843

F-MAMMA1000845//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 (EC 1.6.5.3).//0.43:58:34//DROSOPHI-LA YAKUBA (FRUIT FLY).//P03895

F-MAMMA1000851//CUTICLE COLLAGEN 34.//0.019:107:29//CAENORHABDITIS ELEGANS.//P34687

F-MAMMA1000855//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//0.00098:149:32//HOMO SAPIENS (HUMAN).//Q15428

F-MAMMA1000856//METALLOTHIONEIN (MT).//0.63:39:41//POTAMON POTAMIOS.//P55952

F-MAMMA1000859//GLYCOPROTEIN X PRECURSOR.///0.014:192:28//EQUINE HERPESVIRUS TYPE 1

(STRAIN AB4P) (EHV-1)//P28968
F-MAMMA1000862//DISINTEGRIN KISTRIN (PLATELET AGGREGATION ACTIVATION INHIBITOR.)//1.0:66:

27//AGKISTRODON RHODOSTOMA (MALAYÁN PIT VIPER) (CALLOSELASMA RHODOSTOMA).//P17494
F-MAMMA1000863//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.4e-16:41:68//HOMO SAPIENS (HUMAN).//
P39188

F-MAMMA1000865//SALIVARY PROUNE-RICH PROTEIN II-1 (FRAGMENT).//0.030:100:32//HOMO SAPIENS (HUMAN).//P81489

F-MAMMA1000867//APTOTOXIN IX (PARALYTIC PEPTIDE IX) (PP IX).//0.98:43:32//APTOSTICHUS SCHLIN-GERI (TRAP-DOOR SPIDER).//P49272

F-MAMMA1000875//PROLINE-RICH PEPTIDE P-B.//0.18:21:47//HOMO SAPIENS (HUMAN).//P02814

55 F-MAMMA1000876//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.5e-22:85:71//HOMO SAPIENS (HU-MAN) //P39189

F-MAMMA1000877//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.2e-38:62:74//HOMO SAPIENS (HUMAN).// P39188

- F-MAMMA1000880//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.49:79:32//BOS TAURUS (BOVINE).// P25508
- F-MAMMA1000883//HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN CHROMOSOME X.//0.87:15:60// CAENORHABDITIS ELEGANS.//O11116
- 5 F-MAMMA1000897//INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN H4 PRECURSOR (ITI HEAVY CHAIN H4) (INTER-ALPHA-TRYPSIN INHIBITOR FAMILY HEAVY CHAIN-RELATED PROTEIN) (PLASMA KALLIKREIN SENSITIVE GLYCOPROTEIN 120) (PK-120).//5.3e-17:130.40//HOMO SAPIENS (HUMAN).//Q14624 F-MAMMA1000906
  - F-MAMMA1000905
- 10 F-MAMMA1000908//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//8.0e-17:70:62//HOMO SAPIENS (HU-MAN) //P08547
  - F-MAMMA1000914//HYPOTHETICAL 6.2 KD PROTEIN.//0.97:36:36//THERMOPROTEUS TENAX VIRUS 1 (STRAIN KRA1) (TTV1).//P19299
    F-MAMMA1000991
- 15 F-MAMMA1000931//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.6e-10:49:65//HOMO SAPIENS (HUMAN).//
  - F-MAMMA1000940//MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L32.//0.42:22:54//RECLINOMONAS AMERICANA //021281
  - F-MAMMA1000941//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.3e-25:55:69//HOMO SAPIENS (HUMAN).// P39188
    - F-MAMMA1000942//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.3e-08:36:75//HOMO SAPIENS (HU-MAN).//P39194
  - F-MAMMA1000943

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- F-MAMMA1000956//SMALL HISTIDINE-ALANINE-RICH PROTEIN PRECURSOR (SHARP) (ANTIGEN 57).//
  0.041:122:25//PLASMODIUM FALCIPARUM (ISOLATE FC27 / PAPUA NEW GUINEA).//P04930
  - F-MAMMA1000957//HEAT-STABLE ENTEROTOXIN A2 PRECURSOR (STA2).//0.024:37:37//ESCHERICHIA COLL//Q47185
  - F-MAMMA1000962//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//6.0e-39:61:78//HOMO SAPIENS (HU-MAN).//P39189
- 30 F-MAMMA1000968//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//0.0054:29:72//HOMO SAPIENS (HUMAN).// P39194
  - F-MAMMA1000975//CUTICLE COLLAGEN DPY-2 PRECURSOR.//1.0:93:30//CAENORHABDITIS ELEGANS.// P35799
  - F-MAMMA1000979//PROLINE-RICH PEPTIDE P-B.//0.012:12:66//HOMO SAPIENS (HUMAN),//P02814
    F-MAMMA1000987//HYPOTHETICAL PROTEIN LAMBDA-SP34.//1.0:47:40//MUS MUSCULUS (MOUSE),//
    - F-MAMMA1000998
  - F-MAMMA1001003//PROBABLE E5 PROTEIN.//1.0:52:42//HUMAN PAPILLOMAVIRUS TYPE 33.//P06426
- F-MAMMA1001008//PROGASTRICSIN PRECURSOR (EC 3.4.23.3) (PEPSINOGEN C) (FRAGMENT).//3.2e-14: 131:35//MACACA FUSCATA FUSCATA (JAPANESE MACAQUE).//P03955
  - F-MAMMA1001021//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.016:61:42//STREPTO-MYCES FRADIAE.//P20186
  - F-MAMMA1001024
- F-MAMMA1001030/ILUTROPIN-CHORIOGONADOTROPIC HORMONE RECEPTOR (LH/CG-R) (LSH-R) (LUTEINIZING HOROMINE RECEPTOR) (FRAGMENT)//2.48-20:234:29/GALLUS GALLUS (CHICKEN)// 090674
  - F-MAMMA1001035/!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/1.7e-15:52.78//HOMO SAPIENS (HU-MAN)//P39193 F-MAMMA1001038//NEUROTOXIN II (TOXIN RP-II) (SODIUM CHANNEL TOXIN II) //0.53:25:48//RADIANTHUS
  - F-MAMMA1001038//NEUROTOXIN II (TOXIN RP-II) (SODIUM CHANNEL TOXIN II).//0.53:25:48//RADIANTHU PAUMOTENSIS (SEA ANEMONE) (HETERACTIS PAUMOTENSIS)//P01534
  - F-MAMMA1001041//SPECTRIN BETA CHAIN, ERYTHROCYTE.//6.3e-18:112:43//MUS MUSCULUS (MOUSE).//P15508
    F-MAMMA10011650
  - F-MAMMA1001059//PUTATIVE ATP-DEPENDENT RNA HELICASE C12C2.06.//1.3e-34:187:47//SCHIZOSAC-CHAROMYCES POMBE (FISSION YEAST).//009747
- F-MAMMA1001067//PROTEIN Q300.//0.36:12:75//MUS MUSCULUS (MOUSE).//Q02722
  - F-MAMMA1001073//HEPATOCYTE NUCLEAR FACTOR 3 FORKHEAD HOMOLOG 1 (HFH-1).//1.0:70:37//RAT-TUS NORVEGICUS (RAT).//Q63244

- F-MAMMA1001074//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.00067:163:32//HOMO SAPIENS (HU-MAN) //P08547
- F-MAMMA1001075//RETINOBLASTOMA BINDING PROTEIN 1 (RBBP-1).//0.53:72:34//HOMO SAPIENS (HU-MAN).//P29374
- F-MAMMA1001078//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//5.0e-79:184:73//HOMO SAPIENS (HU-
  - F-MAMMA1001080//IG HEAVY CHAIN PRECURSOR V-III REGION (VH26).//1.7e-27:82:71//HOMO SAPIENS
- (HUMAN).//P01764 F-MAMMA1001082
- 10 F-MAMMA1001091//HYPOTHETICAL BHLF1 PROTEIN.//3.1e-05:198:32//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181
  - F-MAMMA1001092//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//7.1e-21:65:72//HOMO SAPIENS (HU-MAN).//P08547
- F-MAMMA1001105//OVO PROTEIN (SHAVEN BABY PROTEIN).//1.0e-18:68:48//DROSOPHILA MELA-15 NOGASTER (FRUIT FLY).//P51521
  - F-MAMMA1001110//PROCOLLAGEN ALPHA 1(IV) CHAIN PRECURSOR.//0.080:108:37//MUS MUSCULUS (MOUSE).//P02463
    - F-MAMMA1001126//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.3e-07:66:45//HOMO SAPIENS (HU-MAN).//P39189
- F-MAMMA1001133//HYPOTHETICAL 13.2 KD PROTEIN IN RPS4A-BAT2 INTERGENIC REGION.//0.96:43:25// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47174
  - F-MAMMA1001139//HYPOTHETICAL 36.7 KD PROTEIN AH6.2 IN CHROMOSOME II.//5.4e-42:81:62// CAENORHABDITIS ELEGANS.//Q09201
- F-MAMMA1001143//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00014:36:66//HOMO SAPIENS (HUMAN).//
  P39188
  - F-MAMMA1001145

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- F-MAMMA1001154//CSBA PROTEIN.//1.0:39:38//BACILLUS SUBTILIS.//P37953
- F-MAMMA1001161//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/5.2e-23:53:64//HOMO SAPIENS (HUMAN).//
- 30 F-MAMMA1001162//CD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27).//0.69:86:31// MUS MUSCULUS (MOUSE).//P41272
  - F-MAMMA1001181//HYPOTHETICAL 81.0 KD PROTEIN C35D10.4 IN CHROMOSOME III.//0.00010:74:47// CAENORHABDITIS ELEGANS.//Q18486
  - F-MAMMA1001186///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//9.0e-32:44:86//HOMO SAPIENS (HU-MAN).//P39194
    - F-MAMMA1001191/JOCTAMER-BINDING TRANSCRIPTION FACTOR 1 (OTF-1) (NF-A1) (FRAGMENT).//0.096: 40-40/MACROPUS EUGENII (TAMMAR WALLABY).//028466 F-MAMMA1001198//EPIDERMAL GROWTH FACTOR RECEPTOR SUBSTRATE SUBSTRATE 15 (PROTEIN
  - EPS15) (AF-1P PROTEIN).//2.5e-75:204:70//HOMO SAPIENS (HUMAN).//P42566 F-MAMMA1001202//METALLOTHIONEIN-II (MT-II) (MT-IIB).//0.52:46:32//CALLINECTES SAPIDUS
  - (BLUE CRAB),//P55950 F-MAMMA1001203//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!/7.3e-11:82:58//HOMO SAPIENS (HU-
  - MAN).//P39192
    F-MAMMA1001206//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/1.9e-17:67:71//HOMO SAPIENS (HUMAN).//
- - F-MAMMA1001215//9 KD PROTEIN.//1.0:51:33//HOMO SAPIENS (HUMAN).//P13994
  - F-MAMMA1001220//IIII ALU SUBFAMILY SB WARNING ENTRY IIII//3.4e-37:55:87//HOMO SAPIENS (HU-MAN).//P39189
  - F-MAMMA1001222//HYPOTHETICAL 73.6 KD PROTEIN CY49.21.//3.7e-06:168:38//MYCOBACTERIUM TU-BERCULOSIS.//Q10690
  - F-MAMMA1001243
    - F-MAMMA1001244//TRP OPERON LEADER PEPTIDE.//1.0:18:55//SERRATIA MARCESCENS.//P03055
  - F-MAMMA1001249//HYPOTHETICAL 7.2 KD PROTEIN IN RPS2 3'REGION (ORF57).//0.57:23:34//ASTASIA LONGA (EUGLENOPHYCEAN ALGA).//P34774
- 55 F-MAMMA1001256//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.3e-07:79:44//HOMO SAPIENS (HUMAN).// P39188
  - F-MAMMA1001259//PUTATIVE DNA HELICASE II HOMOLOG (EC 3.6.1.-).//0.046:86:32//MYCOPLASMA GEN-ITALIUM.//P47486

- F-MAMMA1001260//MYOSIN HEAVY CHAIN, PERINATAL SKELETAL MUSCLE.//2.7e-05:219:27//HOMO SAPIENS (HUMAN).//P13535
- F-MAMMA1001268//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//9.7e-27:89:67//HOMO SAPIENS (HU-MAN).//P08547
- F-MAMMA1001271//ATROPHIN-1 (DENTATORUBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN).//4.0e-06:126: 38//HOMO SAPIENS (HUMAN).//P54259
  - F-MAMMA1001274/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.4e-29:57:66//HOMO SAPIENS (HU-MAN) //P39194
- F-MAMMA1001280//BACTERIOCIN MICROCIN B17 PRECURSOR (MCB17).//0.27:24:54//ESCHERICHIA CO-10 LL//P05834
  - F-MAMMA1001292//HYPOTHETICAL PROTEIN KIAA0176 (FRAGMENT).//1.3e-73:208:69//HOMO SAPIENS (HUMAN).//Q14681
  - F-MAMMA1001296//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//6.9e-22:41:80//HOMO SAPIENS (HU-MAN) //P39193
- F-MAMMA1001298//HYPOTHETICAL PROTEIN HI0371.//0.99:29:37//HAEMOPHILUS INFLUENZAE.//P44668 F-MAMMA1001305//GTPASE-ACTIVATING PROTEIN RHOGAP (RHO-RELATED SMALL GTPASE PROTEIN ACTIVATOR) (CDC42 GTPASE-ACTIVATING PROTEIN) (P50-RHOGAP).//9.99-62:222:54//HOMO SAPIENS (HUMAN).//Q07960
  - F-MAMMA1001322//B-CELL GROWTH FACTOR PRECURSOR (BCGF-12 KD).//2.1e-09:46:60//HOMO SAPI-ENS (HUMAN).//P20931
    - F-MAMMA1001324//POL POLYPROTEIN [CONTAINS: PROTEASE (EC 3.4.23.-); REVERSE TRANSCRIPTASE (EC 2.7.7.49); RIBONUCLEASE H (EC 3.1.26.4)].//2.56-43:128:50//FRIEND MURINE LEUKEMIA VIRUS (ISO-LATE PVC-211) (F-MULV)./P2808
- F-MAMMA1001330//HEMOGLOBIN ZETA CHAIN (FRAGMENTS).//0.30:51:37//MACROPUS EUGENII (TAM-MAR WALLABY).//P81044
  - F-MAMMA1001341//TRISTETRAPROLINE (TTP) (TIS11A) (TIS11) (ZFP-36) (GROWTH FACTOR- INDUCIBLE NUCLEAR PROTEIN NUP475).//0.024:89:39//HOMO SAPIENS (HUMAN).//P26651
  - F-MAMMA1001343//PROBABLE E5 PROTEIN.//0.60:64:29//HUMAN PAPILLOMAVIRUS TYPE 16.//P06927 F-MAMMA1001346//PROTEINASE INHIBITOR IIB (FRAGMENTS).//0.97:33:45//SOLANUM TUBEROSUM (PO-
- 30 TATO).//P01082 F-MAMMA1001383//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/1.2e-30:86:77//HOMO SAPIENS (HU-
  - MAN).//P39194
    F-MAMMA1001388//LEUCINE-RICH ALPHA-2-GLYCOPROTEIN (LRG).//9.2e-91:195:92//HOMO SAPIENS
    (HUMAN).//P02750
  - F-MAMMA1001397///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.5e-19:55:69//HOMO SAPIENS (HUMAN).//
    - F-MAMMA1001408//SALIVARY GLUE PROTEIN SGS-7 PRECURSOR.//0.60:45:35//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P02841
- F-MAMMA1011411/IGLUCOAMYLASE S1/S2 PRECURSOR (EC 3.2.1.3) (GLUCAN 1.4-ALPHA- GLUCOSI-DASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//5.8e-06.153:26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P08640
  - F-MAMMA1001419//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.3e-16:99:51//HOMO SAPIENS (HU-MAN) //P39104
- F-MAMMA1001420//!!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!!//0.0018:23:65//HOMO SAPIENS (HU-MAN) //P39190
  - F-MAMMA1001435//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.7e-22:60:58//HOMO SAPIENS (HU-MAN).//P39195
    - F-MAMMA1001442

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- F-MAMMA1001452//GENE 35 PROTEIN (GP35).//0.61:31:45//MYCOBACTERIOPHAGE L5.//Q05245 F-MAMMA1001465//HYPOTHETICAL PROTEIN E-115.//0.0026:68:38//HUMAN ADENOVIRUS TYPE 2.// P03990
- F-MAMMA1001476//URIDINE KINASE (EC 2.7.1.48) (URIDINE MONOPHOSPHOKINASE) (FRAGMENT).//
  3.7e-94:201:92//MUS MUSCULUS (MOUSE).//P52623
- F-MAMMA1001487//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.6e-16:89:41//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548
  - F-MAMMA1001501//CALPAIN 1, LARGE [CATALYTIC] SUBUNIT (EC 3.4.22.17) (CALCIUM-ACTIVATED NEU-

- TRAL PROTEINASE) (CANP) (MU-TYPE).//6.2e-59:86:97//HOMO SAPIENS (HUMAN).//P07384
- F-MAMMA1001502//HYPOTHETICAL 11.4 KD PROTEIN (ORF1).//0.21:79:30//STREPTOMYCES FRADIAE.//

### F-MAMMA1001510

- F-MAMMA1001522//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.67:98:31//STREPTO-MYCES FRADIAE //P20186
  - F-MAMMA1001547//PROBABLE MOLYBDENUM-PTERIN BINDING PROTEIN.//0.97:35:42//HAEMOPHILUS
  - INFLUENZAE.//P45183
    F-MAMMA1001551//HYPOTHETICAL PROTEIN MJ0458.1.//0.038:31:41//METHANOCOCCUS JANNASCHII.//
- 10 P81308

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- F-MAMMA1001576
  F-MAMMA1001576/JTUBULIN GAMMA CHAIN.//1.6e-86:162:99//XENOPUS LAEVIS (AFRICAN CLAWED EPOC).//293330
- F-MAMMA1001590//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.0035;38:55//HOMO SAPIENS (HUMAN).//
- F-MAMMA1001600//CONNECTIVE TISSUE GROWTH FACTOR PRECURSOR.//0.85:53:33//HOMO SAPIENS
  - F-MAMMA1001604//HYPOTHETICAL 11.1 KD PROTEIN C30D11.02C IN CHROMOSOME I.//0.14:82:29// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//009902
- F-MAMMA1001606//HIGH MOBILITY GROUP PROTEIN HMGI-C.//8.2e-05:77:37//HOMO SAPIENS (HUMAN).//
- F-MAMMA1001620///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//5.5e-05:24:66//HOMO SAPIENS (HU-
- F-MAMMA1001627//CUTICLE COLLAGEN 40.//0.82:131:31//CAENORHABDITIS ELEGANS.//P34804
- 25 F-MAMMA1001630//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//8.6e-26:57:78//HOMO SAPIENS (HU-MAN).//P39194
  - F-MAMMA1001633//ZINC FINGER PROTEIN 165.//6.9e-38:160:55//HOMO SAPIENS (HUMAN).//P49910 F-MAMMA1001635
- F-MAMMA1001649//SPERM PROTAMINE P1.//0.39:31:41//TACHYGLOSSUS ACULEATUS ACULEATUS (AUS-™ TRALIAN ECHIDNA).//P35311
- F-MAMMA1001654//NON-RECEPTOR TYROSINE KINASE SPORE LYSIS A (EC 2.7.1.112) (TYROSINE- PRO-TEIN KINASE 1).//5.6e-06:99:28//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P18160
  - F-MAMMA1001663//VERY HYPOTHETICAL XYLU PROTEIN.//0.99:27:37//ESCHERICHIA COLI.//P05056
  - F-MAMMA1001670//CUTICLE COLLAGEN 1.//0.033:97:37//CAENORHABDITIS ELEGANS.//P08124
- 35 F-MAMMA1001671
  - F-MAMMA1001679//PROCOLLAGEN ALPHA 2(IV) CHAIN PRECURSOR.//0.92:32:50//HOMO SAPIENS (HU-MAN).//P08572
  - F-MAMMA1001683//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.00026:147:34//STREP-TOMYCES FRADIAE.//P20186
- 40 F-MAMMA1001686
  - F-MAMMA1001692//SMALL HYDROPHOBIC PROTEIN (SMALL PROTEIN 1A), I/1.0:34:26//BOVINE RESPIRA-TORY SYNCYTIAL VIRUS (STRAIN A51908) (BRS), I/P24616
  - F-MAMMA1001711//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//1.1e-28:56:69//HOMO SAPIENS (HU-MAN) //P39194
- 45 F-MAMMA1001715//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.6e-08:39:71//HOMO SAPIENS (HUMAN).//
  - F-MAMMA1001730//METALLOTHIONEIN-B (MTB).//1.0:17:64//STRONGYLOCENTROTUS PURPURATUS (PURPLE SEA URCHIN).//Q27287
  - F-MAMMA1001735//TUBULIN BETA-5 CHAIN (CLASS-V).//5.1e-121:213:97//GALLUS GALLUS (CHICKEN).// P09653
  - F-MAMMA1001740

- F-MAMMA1001743//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.3e-09:100:42//HOMO SAPIENS (HU-MAN).//P39195
- F-MAMMA1001744//POU DOMAIN PROTEIN 2.//0.97:59:38//BRACHYDANIO RERIO (ZEBRAFISH) (ZEBRA
  55 DANIO).//Q90270
- F-MAMMA1001745//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//2.1e-43:199:42//HOMO SAPIENS (HU-MAN).//P08547
  - F-MAMMA1001751//TWK-8 PROTEIN.//2.9e-15:77:36//CAENORHABDITIS ELEGANS.//P34410

- F-MAMMA1001754//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.019:20:45//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01645
- F-MAMMA1001757//HYPOTHETICAL 9.2 KD PROTEIN IN RNPA 3'REGION.//0.94:30:43//PSEUDOMONAS PUTIDA.//P25753
- F-MAMMA1001760///!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!/4.6e-34:103:59//HOMO SAPIENS (HU-MAN).//P39191
  - F-MAMMA1001764
  - F-MAMMA1001768//HYPOTHETICAL PROTEIN UL61.//0.042:167:33//HUMAN CYTOMEGALOVIRUS (STRAIN AD169).//P16818
- 10 F-MAMMA1001769/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.0e-29:97:69//HOMO SAPIENS (HU-MAN) //P39194
  - F-MAMMA1001771//TRANSMEMBRANE PROTEIN SEX PRECURSOR.//3.3e-09:123:32//HOMO SAPIENS (HUMAN).//P51805
  - F-MAMMA1001783///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-09:55:61//HOMO SAPIENS (HUMAN).//
- F-MAMMA1001785//RAS-RELATED PROTEIN RABC.//1.9e-06:120:25//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P34143
  - F-MAMMA1001788//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//9.3e-29:46:76//HOMO SAPIENS (HU-MAN).//P08547
- 20 F-MAMMA1001790//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.7e-24:69:69//HOMO SAPIENS (HUMAN).// P39188
  - F-MAMMA1001806//HYPOTHETICAL 21.2 KD PROTEIN IN TOR2-MNN4 INTERGENIC REGION.//0.95:58:36// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36042
- F-MAMMA1001812//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//8.8e-12:53:69//HOMO SAPIENS (HU-MAN).//P39195
  - F-MAMMA1001815//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.11:30:70//HOMO SAPIENS (HU-MAN).//P08547
  - F-MAMMA1001817//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.9e-16:86:55//HOMO SAPIENS (HUMAN).// P39188
- 30 F-MAMMA1001818

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- F-MAMMA1001820//VITTELLINE MEMBRANE PROTEIN VM26AB PRECURSOR (PROTEIN TU-4) (PROTEIN SV23).//0.0030:63:42//DROSOPHILA MELANOGASTER (FRUIT FLY).//P13238
  - F-MAMMA1001824//APTOTOXIN VII (PARALYTIC PEPTIDE VII) (PP VII).//0.99:26:34//APTOSTICHUS SCHLIN-GERI (TRAP-DOOR SPIDER).//P49271
- 35 F-MAMMA1001836/!!!! ALU SUBFAMILY SX WARNING ENTRY!!!!//2.6e-35:77:88//HOMO SAPIENS (HU-MAN).//P39195
  - F-MAMMA1001837//ZINC FINGER PROTEIN 191.//1.3e-27:106:58//HOMO SAPIENS (HUMAN).//O14754 F-MAMMA1001848//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.0e-19:92:58//HOMO SAPIENS (HUMAN).// P39188
- 40 F-MAMMA1001851

- F-MAMMA1001854
  - F-MAMMA1001858//ISOTOCIN-NEUROPHYSIN IT 1 PRECURSOR.//0.93:42:38//CATOSTOMUS COMMERSO-NI (WHITE SUCKER).//P15210
- F-MAMMA1001864//PROBABLE ABC TRANSPORTER PERMEASE PROTEIN MG189.//0.77:161:27//MYCO-PLASMA GENITALIUM.//P47435
  - F-MAMMA1001888//FK508-BINDING NUCLEAR PROTEIN (PEPTIDYL-PROLYL CIS-TRANS ISOMERASE) (PPIASE) (EC 5.2.1.8) (PROLINE ROTAMASE) (NUCLEOLAR PROLINE ISOMERASE) (FKBP-70).//0.00013: 219:26//SACCHAROMYCES CEREVISIAE (BAKERS YEAST)//P38911
  - F-MAMMA1001874//SPERM HISTONE P2 PRECURSOR (PROTAMINE MP2).//0.0075:76:31//MUS MUSCULUS (MOUSE).//P07978
    - F-MAMMA1001878//GLYCINE-RICH CELL WALL STRUCTURAL PROTEIN (CLONE W10-1) (FRAGMENT).// 0.020:10:80//LYCOPERSICON ESCULENTUM (TOMATO).//Q01157
    - F-MAMMA1001880
- F-MAMMA1001890//!!!! ALU SUBFAMILY SC WARNING ENTRY!!!!//5.1e-34:56:83//HOMO SAPIENS (HU-
  - F-MAMMA1001907//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.7e-12:44:68//HOMO SAPIENS (HU-MAN).//P39194
  - F-MAMMA1001908//HYPOTHETICAL 16.2 KD PROTEIN IN PRP24-RRN9 INTERGENIC REGION.//0.00013:77:

37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q03525

F-MAMMA1001931//HYPOTHETICAL 118.2 KD PROTEIN F43C1.1 IN CHROMOSOME III.//0.41:106:29// CAENORHABDITIS ELEGANS.//Q09564

F-MAMMA1001956//OCTAPEPTIDE-REPEAT PROTEIN T2://0.00053:149:30//MUS MUSCULUS (MOUSE).//

F-MAMMA1001963//HYPOTHETICAL PROTEIN IN NAC 5'REGION (ORF X) (FRAGMENT).//1.0:46:28//KLEB-SIELLA AEROGENIES //OR8600

F-MAMMA1001969//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.7e-34:97:68//HOMO SAPIENS (HU-MAN).//P08547

10 F-MAMMA1001970//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.2e-07:67:37//HOMO SAPIENS (HU-MAN).//P08547

F-MAMMA1001992//PROTEIN Q300.//0.53:14:71//MUS MUSCULUS (MOUSE).//Q02722

F-MAMMA1002009//PROBABLE E5 PROTEIN.//0.17:56:32//HUMAN PAPILLOMAVIRUS TYPE 31.//P17385 F-MAMMA1002011//MYRISTOYLATED ALANINE-RICH C-KINASE SUBSTRATE (MARCKS) (PROTEIN KINASE

15 C SUBSTRATE, 80 KD PROTEIN, LIGHT CHAIN) (PKCSL) (80K-L PROTEIN).//1.0:100:31//HOMO SAPIENS (HUMAN).//P29966

F-MAMMA1002032//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.1e-21:86:65//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1002033//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//8.5e-20:67:58//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1002041//MALE SPECIFIC SPERM PROTEIN MST84DC.//1.0:17:52//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01644

F-MAMMA1002042//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//0.19:45:46//HOMO SAPIENS (HUMAN).// P39192

F-MAMMA1002047//TYROSINE AMINOTRANSFERASE (EC 2.6.1.5) (L-TYROSINE:2-OXOGLUTARATE AMI-NOTRANSFERASE) (TAT).//0.0017:50:46//RATTUS NORVEGICUS (RAT).//P04694

F-MAMMA1002056//iiii ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.2e-37:70:77//HOMO SAPIENS (HU-MAN).//P39194

F-MAMMA1002058//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-08:26:76//HOMO SAPIENS (HUMAN).//

F-MAMMA1002068//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//2.0e-11:78:46//HOMO SAPIENS (HU-MAN).//P08547

F-MAMMA1002078//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.96:26:46//COTURNIX COTURNIX JAPONICA (JAPANESE QUAIL).//P50682

F-MAMMA1002082//SUPPRESSOR PROTEIN SRP40.//0.23:95:32//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583

F-MAMMA1002084//HYPOTHETICAL 7.5 KD PROTEIN.//1.0:40:35//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20520

F-MAMMA1002093

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40 F-MAMMA1002108//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.00079:143:33//STREP-TOMYCES FRADIAF //P20186

F-MAMMA1002118//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:43:34//METRIDIUM SENILE (BROWN SEA ANEMONE) (FRILLED SEA ANEMONE).//O47493

F-MAMMA1002125//!!!! ALU SUBFAMILY SC WARNING ENTRY!!!!//1.9e-14:60:68//HOMO SAPIENS (HU-MAN).//P39192

F-MAMMA1002132

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F-MAMMA1002140//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.4e-24:69:65//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1002143//SERUM PROTEIN MSE55.//2.1e-16:166:43//HOMO SAPIENS (HUMAN).//Q00587

F-MAMMA1002145//36.4 KD PROLINE-RICH PROTEIN.//0.00014:84:29//LYCOPERSICON ESCULENTUM (TO-MATO).//Q00451

F-MAMMA1002153

F-MAMMA1002155

F-MAMMA1002156//METALLOPROTEINASE INHIBITOR PRECURSOR.//0.90:58:34//STREPTOMYCES NI-GRESCENS.//P01077

F-MAMMA1002158

F-MAMMA1002170//40S RIBOSOMAL PROTEIN S2 (S4) (LLREP3 PROTEIN).//6.0e-66:157:70//HOMO SAPIENS (HUMAN).//P15880

- F-MAMMA1002174//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//6.5e-25:56:64//HOMO SAPIENS (HUMAN).//
- F-MAMMA1002198//THIOREDOXIN PEROXIDASE 1 (THIOREDOXIN-DEPENDENT PEROXIDE REDUCTASE 1) (THIOL-SPECIFIC ANTIOXIDANT PROTEIN) (TSA) (PRP) (NATURAL KILLER CELL ENHANCING FACTOR
- B) (NKEF-B)//8, 06-09:28:100/InOMO SAPIENS (HUMAN)//P32119 F-MAMMA1002209/ITRANSCRIPTION INTIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII135) (TAFII-130) (TAFII130)//0.0023:132:33/I+OMO SAPIENS (HUMAN)//000268
  - F-MAMMA1002215//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//0.00032:68:35//HOMO SAPIENS (HU-MAN).//P02452
- 10 F-MAMMA1002219//INTRACELLULAR PROTEIN TRANSPORT PROTEIN USO1.//0.0079:224:24//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P25386
  - F-MAMMA1002230

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- F-MAMMA1002236//TRANSLATION INITIATION FACTOR EIF-2B GAMMA SUBUNIT (EIF-2B GDP-GTP EX-CHANGE FACTOR).//1.4e-118:151:94//RATTUS NORVEGICUS (RAT).//P70541
- 15 F-MAMMA1002243//WISKOTT-ALDRICH SYNDROME PROTEIN HOMOLOG (WASP).//0.028:112:33//MUS MUSCULUS (MOUSE).//P70315
  - F-MAMMA1002250//T-CELL RECEPTOR BETA CHAIN PRECURSOR (ANA 11).//0.0012:80:32//ORYCTOLA-GUS CUNICULUS (RABBIT).//P06333
  - F-MAMMA1002267//ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6).//0.17:139:28//TRYPANOSOMA BRUCFI BRUCFI //P24499
  - F-MAMMA1002268//60S RIBOSOMAL PROTEIN L22.//0.00026:163:30//DROSOPHILA MELANOGASTER (FRUIT FLY).///750887
  - F-MAMMA1002269//HISTIDINE-RICH, METAL BINDING POLYPEPTIDE.//0.35:14:57//HELICOBACTER PY-LORI (CAMPYLOBACTER PYLORI).//Q48251
- 25 F-MAMMA1002282/!/!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//6.1e-05:32:65//HOMO SAPIENS (HU-MAN).//P39192
  - F-MAMMA1002292//TROPOMYOSIN 2.//1.4e-05:100:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40414
  - F-MAMMA1002293///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//6.8e-25:127:44//HOMO SAPIENS (HUMAN).//
- F-MAMMA1002294//ALPHA TRANS-INDUCING PROTEIN (ALPHA-TIF).//0.00011:138:38//BOVINE HERPESVI-RUS TYPE 1 (STRAIN P8-2).//P30020
  - F-MAMMA1002297/IA-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.15:144:30//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323
- 35 F-MAMMA1002298//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//1.0e-05:40:50//MUS MUSCULUS (MOUSE).//P05143
  - F-MAMMA1002299//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3),//0.84:65:32//STRUTHIO CAMELUS (OSTRICH),//O21405
- F-MAMMA1002308//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.3e-29:61:73//HOMO SAPIENS (HUMAN).//
  40 P39188
- F-MAMMA1002310//SPERM MITOCHONDRIAL CAPSULE SELENOPROTEIN (MCS).//0.00016:70:38//MUS MUSCULUS (MOUSE).//P15265
- F-MAMMA1002311//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//9.4e-09:84:54//HOMO SAPIENS (HU-
- 45 F-MAMMA1002312//HYPOTHETICAL 10.8 KD PROTEIN IN GP30-RIII INTERGENIC REGION (URF Y).//0.48: 48:33//BACTERIOPHAGE 14.//P33084
- F-MAMMA1002317
  F-MAMMA1002319//RETROVIRUS-RELATED POL POLYPROTEIN ICONTAINS: REVERSE TRANSCRIPTASE
  - (EC 2.7.7.49); ENDONUCLEASEJ,//0.011:128:27//MUS MUSCULUS (MOUSE),//P11389
    F-MAMMA1002322//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/f.2e-20:92:57//HOMO SAPIENS (HU-
  - MAN), I/P39195

    F-MAMMA1002329//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L), I/O.051:33:36//XENOPUS LAEVIS (AFRI-
    - CAN CLAWED FROG).//P03931
- F-MAMMA1002332//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.5e-20:116:51//HOMO SAPIENS (HU-55 MAN).//P08547
  - F-MAMMA1002333//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//0.0017:214:31//BOS TAURUS (BO-VINE).//P02453
  - F-MAMMA1002339//COPPER-METALLOTHIONEIN (CU-MT).//0.59:42:38//HELIX POMATIA (ROMAN SNAIL)

(EDIBLE SNAIL) //P55947

F-MAMMA1002347//!!!! ALU SUBFAMILY J WARNING ENTRY!!!!/0.43:26:61//HOMO SAPIENS (HUMAN).//

F-MAMMA1002351//HYPOTHETICAL PROTEIN MJ0304.//2.3e-07:139:25//METHANOCOCCUS JANNAS-CHII //Q57752

F-MAMMA1002352

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F-MAMMA1002353//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00028:31:80//HOMO SAPIENS (HUMAN).//

F-MAMMA1002355//!!!! ALU SUBFAMILY SP WARNING ENTRY!!!!//4.2e-28:87:73//HOMO SAPIENS (HU-MAN).//739193

MAN).//P39193 F-MAMMA1002356//RELAXIN.//0.95:31:35//SQUALUS ACANTHIAS (SPINY DOGFISH).//P11953

F-MAMMA1002359//CHLOROPLAST 50S RIBOSOMAL PROTEIN L33.//0.93:44:36//GUILLARDIA THETA (CRYPTOMONAS PHI).//0.78487

F-MAMMA1002360/LATE L2 MU CORE PROTEIN PRECURSOR (PROTEIN X).//0.94:30:43//BOVINE ADENO-VIRUS TYPE 2 (MASTADENOVIRUS BOS2).//Q96626

F-MAMMA1002361//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.0e-08:45:68//HOMO SAPIENS (HUMAN).//

F-MAMMA1002362//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.58:23:26//LUMBRICUS TERRESTRIS (COMMON EARTHWORM).//034942

F-MAMMA1002380//SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.23:100:27//DROSOPHILA SIMU-LANS (FRUIT FLY).//P13729

F-MAMMA1002384

F-MAMMA1002385//HYPOTHETICAL 40.9 KD PROTEIN IN ORC2-TIP1 INTERGENIC REGION.//3.8e-14:125: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38241

25 F-MAMMA1002392//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:17:58//BRANCHIOSTOMA LANCEO-LATUM (COMMON LANCELET) (AMPHIOXUS).//021003

F-MAMMA1002411//30S RIBOSOMAL PROTEIN \$17.//0.85:49:32//SYNECHOCYSTIS \$P. (STRAIN PCC 6803).//P73311

F-MAMMA1002413//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 1 (EC 1.6.5.3) (FRAGMENT).//0.97:41: 39//DROSOPHILA AFFINIS (FRUIT FLY).//P51926

F-MAMMA1002417//RFBJ PROTEIN.//0.99:31:35//SHIGELLA FLEXNERI.//P37786

F-MAMMA1002427//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//1.6e-33:135:59//HOMO SAPIENS (HU-MAN).//P39194

F-MAMMA1002428//HYPOTHETICAL PROTEIN C18.//0.97.34:44//SWINEPOX VIRUS (STRAIN KASZA) (SPV).//P32217

F-MAMMA1002434//!!! ALU SUBFAMILY SB WARNING ENTRY!!!!/3.1e-36:56:78//HOMO SAPIENS (HU-

F-MAMMA1002434//!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//3.1e-36:56:78//HOMO SAPIENS MAN).//P39189

F-MAMMA1002446

F-MAMMA1002454//EARLY NODULIN 20 PRECURSOR (N-20).//0.77:57:45//MEDICAGO TRUNCATULA (BAR-REL MEDIC).//P93329

F-MAMMA1002461//VASODILATOR-STIMULATED PHOSPHOPROTEIN (VASP).//1.3e-05:193:32//CANIS FA-MILIARIS (DOG).//P50551

F-MAMMA1002470//HYPOTHETICAL 80.7 KD PROTEIN IN ERG7-NMD2 INTERGENIC REGION.//1.0e-75:231: 60//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38795

45 F-MAMMA1002475//POSSIBLE GLOBAL TRANSCRIPTION ACTIVATOR SNF2L4 (SNF2-BETA) (BRG-1 PRO-TEIN) (MITOTIC GROWTH AND TRANSCRIPTION ACTIVATOR) (BRAHMA PROTEIN HOMOLOG 1).//0.013: 99:30//HOMO SAPIENS (HUMAN).//P51532

F-MAMMA1002480//NONSTRUCTURAL PROTEIN 5B.//1.0:23:43//HUMAN CORONAVIRUS (STRAIN 229E).// P19741

F-MAMMA1002485//STANNIOCALCIN PRECURSOR.//2.1e-23:88:46//HOMO SAPIENS (HUMAN).//P52823 F-MAMMA1002494//MOLT-INHIBITING HORMONE (MIH).//1.0:32:37//PROCAMBARUS CLARKII (RED SWAMP CRAYFISH).//P58848

F-MAMMA1002498//6.7 KD PROTEIN (ORF 5).//1.0:26:42//BARLEY YELLOW DWARF VIRUS (ISOLATE PAV) (BYDV).//P09517

F-MAMMA1002524/HYPOTHETICAL 117.8 KD PROTEIN IN STE2-FRS2 INTERGENIC REGION.//5.0e-26:222: 35/SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43571

F-MAMMA1002530//CYTOSOLIC PHOSPHOLIPASE A2 (EC 3.1.1.4) (CPLA2) (PHOSPHATIDYLCHOLINE 2-ACYLHYDROLASE) / LYSOPHOSPHOLIPASE (EC 3.1.1.5).//4.5e-12:88:44//HOMO SAPIENS (HUMAN).//

### P47712

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F-MAMMA1002545//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//4.3e-29:97:71//HOMO SAPIENS (HU-MAN) //P39195

F-MAMMA1002554//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RPB1) (FRAG-MENT).//0.46:54:40//CRICETULUS GRISEUS (CHINESE HAMSTER).//P11414

F-MAMMA1002556//METALLOTHIONEIN 20-I ISOFORMS A AND B (MT-20-IA AND MT-20-IB).//0.99:21:47// MYTH US EDULIS (BLUE MUSSEL).//P80251

F-MAMMA1002566//TRANSCRIPTION FACTOR P65 (NUCLEAR FACTOR NF-KAPPA-B P65 SUBUNIT).//0.70: 130:30//MUS MUSCULUS (MOUSE).//Q04207

10 F-MAMMA1002571//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (FRAGMENT).// 0.54:45:51//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P35084

F-MAMMA1002573//PARATHYMOSIN.//1.5e-07:69:46//HOMO SAPIENS (HUMAN).//P20962

F-MAMMA1002585//MYOSIN LIGHT CHAIN 1, SLOW-TWITCH MUSCLE B/VENTRICULAR ISOFORM (FRAG-MENT).//0.38:36:36//MUS MUSCULUS (MOUSE).//P09542

15 F-MAMMA1002590//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.99:22:77//HOMO SAPIENS (HUMAN).// P39195

F-MAMMA1002597/!!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/2.1e-18:44:70//HOMO SAPIENS (HU-MAN).//P39194

F-MAMMA1002598//60S RIBOSOMAL PROTEIN L7.//1.8e-16:40:100//HOMO SAPIENS (HUMAN).//P18124 F-MAMMA1002603

F-MAMMA1002612//30S RIBOSOMAL PROTEIN S16 (FRAGMENT).//1.0:29:37//THERMUS AQUATICUS.//
O07348

F-MAMMA1002617//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT).// 0.00041:81:34//RATTUS NORVEGICUS (RAT).//P10164

25 F-MAMMA1002618//ESCARGOT/SNAIL PROTEIN HOMOLOG (FRAGMENT).//0.11:18:50//PSYCHODA CINE-REA.//Q02027

F-MAMMA1002619/PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE K02C4.3 (EC 3.1.2.15) (UBIQUITIN THIOLESTERASE) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE) (DEUBIQUITINATING EN-ZYME) //1.8e-13.110-40/CAENORHABDITIS ELEGANS //0.09931

F-MAMMA1002622//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/8.4e-05:53:58//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1002823/I/PEPTIDYL-GLYCINE ALPHA-AMIDATING MONOOXYGENASE PRECURSOR (EC 1.14.17.3) (PAM)/I/2-6-07:37:78/I/HOMO SAPIENS (HUMAN) //P19021 F-MAMMA1002825

F-MAMMA1002629//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.4e-19:49:73//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1002636//COLLAGEN ALPHA 2(VI) CHAIN (FRAGMENT).//1.7e-07:189:32//HOMO SAPIENS (HU-MAN).//P12110

F-MAMMA1002637/IKINESIN LIGHT CHAIN (KLC)//7.7e-54:227:52/IRATTUS NORVEGICUS (RAT).//P37285
40 F-MAMMA1002646/INEUROFILAMENT TRIPLET H PROTEIN (200 KD NEUROFILAMENT PROTEIN) (NF-H).//
0.034:199:25/I/MUS MUSCULUS (MOUSE)://P19246

F-MAMMA1002650//TRANSCRIPTION REGULATOR PROTEIN BACH2 (BTB AND CNC HOMOLOG 2).//1.7e-07:104:32//MUS MUSCULUS (MOUSE).//P97303

F-MAMMA1002655//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//1.0:25:44//HOMO SAPIENS (HUMAN).//P22532

F-MAMMA1002662

F-MAMMA1002662
F-MAMMA1002665/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/2.3e-07:54:57//HOMO SAPIENS (HU-

F-MAMMA1002671//ACETYL-COENZYME A SYNTHETASE (EC 6.2.1.1) (ACETATE--COA LIGASE) (ACYL- AC-TIVATING ENZYME).//1.4e-10:144:31//ESCHERICHIA COLI.//P27550

F-MAMMA1002673//BREVICAN CORE PROTEIN PRECURSOR.//0.76:64:39//BOS TAURUS (BOVINE).//

F-MAMMA1002684//HYPOTHETICAL 11.8 KD PROTEIN IN GP55-NRDG INTERGENIC REGION.//0.094:77: 27/BACTERIOPHAGE T4.//P07079

55 F-MAMMA1002685//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//0.0017:177:34//RATTUS NORVEGICUS (RAT).//P02454

F-MAMMA1002698

F-MAMMA1002699//HYPOTHETICAL 45.1 KD PROTEIN IN RPS5-ZMS1 INTERGENIC REGION.//1,2e-28:127:

- 47//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47160
- F-MAMMA1002701//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.0:14:92//HOMO SAPIENS (HUMAN).//
- F-MAMMA1002708//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//7.9e-27:52:65//HOMO SAPIENS (HU-MAN).//P39193
  - F-MAMMA1002711//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.7e-24:54:75//HOMO SAPIENS (HUMAN).// P39188

### F-MAMMA1002721

- F-MAMMA1002727//SOX-13 PROTEIN (FRAGMENT).//0.70:36:38//MUS MUSCULUS (MOUSE).//Q04891

  F-MAMMA1002728//HYPOTHETICAL 6.0 KD PROTEIN.//1.0:25:44//THERMOPROTEUS TENAX VIRUS 1
  - (STRAIN KRA1) (TTV1)//P19305
    F-MAMMA1002744//HYPOTHETICAL 13.4 KD PROTEIN IN ACT5-YCK1 INTERGENIC REGION.//1.0:52:34//
    SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P38834
- F-MAMMA1002746//HYPOTHETICAL 5.6 KD PROTEIN (ORF A-45).//1.0:22:40//SULFOLOBUS VIRUS-LIKE PARTICLE SSV1.//P20198

## F-MAMMA1002748

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- F-MAMMA1002754//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.1e-21:56:64//HOMO SAPIENS (HUMAN).//
- F-MAMMA1002758//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.37:14:64//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01645
- F-MAMMA1002764//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//4.7e-32:79:60//HOMO SAPIENS (HU-MAN)//P39194

### F-MAMMA1002765//PARATHYMOSIN.//0.79:63:28//BOS TAURUS (BOVINE).//P08814

- F-MAMMA1002769/IGAR2 PROTEIN.//0.00037:192:27//SCHIZOSACCHAROMYCES POMBE (FISSION VEAST) ///P41801
- F-MAMMA1002775//HYPOTHETICAL 36.7 KD PROTEIN C2F7.14C IN CHROMOSOME I.//5.4e-54:240:49// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09704
  - F-MAMMA1002780
    F-MAMMA1002782//MARGATOXIN (MGTX).//1.0:31:38//CENTRUROIDES MARGARITATUS (SCORPION).//
- P40755 F-MAMMA1002796///CE NUCLEATION PROTEIN //0.0018:100:41//PSEUDOMONAS FLUORESCENS //
  - P09815
    F-MAMMA1002807//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.3e-23:100:59//HOMO SAPIENS (HUMAN).//
  - P39188
    F-MAMMA1002820//NEUROTOXIN IV (LQQ IV).//1.0:18:50//LEIURUS QUINQUESTRIATUS QUINQUESTRIA-
    - TUS (EGYPTIAN SCORPION).//P01489
      F-MAMMA1002830////II ALU SUBFAMILY SX WARNING ENTRY IIII//4.7e-24:55:74//HOMO SAPIENS (HU-MAN) //P39195
    - F-MAMMA1002833//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.6e-31:95:73//HOMO SAPIENS (HU-MAN).//P39189
  - F-MAMMA1002835//HYPOTHETICAL 42.1 KD PROTEIN F13G3.3 IN CHROMOSOME 1.//1.0:54:37//
    CAENORHABDITIS ELEGANS.//O19417
    - F-MAMMA1002838//!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.5e-27:99:70//HOMO SAPIENS (HU-MAN) //P39193
- 45 F-MAMMA1002842//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.3e-13:65:63//HOMO SAPIENS (HU-MAN).//P39195
  - F-MAMMA1002843//METALLOTHIONEIN-II (MT-II).//0.97:19:47//MUS MUSCULUS (MOUSE).//P02798
  - F-MAMMA100284/IHYPOTHETICAL 24 1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION./I4.9e-08:119:36// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV)./IP41479 F-MAMMA1002858/IATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (AGL)/I0.98:37:37:I/PAN TROGLODYTES
  - (CHIMPANZEE).//Q35647 F-MAMMA1002868//III ALU SUBFAMILY J WARNING ENTRY !!!//3.8e-10:51:62//HOMO SAPIENS (HUMAN).//
    - P39188
      F-MAMMA1002869//PINCH PROTEIN (PARTICULARY INTERESTING NEW CYS-HIS PROTEIN).//1.8e-95:194:
- 78//HOMO SAPIENS (HUMAN).//P48059
  F-MAMMA1002871//G-PROTEIN COUPLED RECEPTOR HOMOLOG R33.//1.0:51:35//RAT CYTOMEGALOVI-
  - RUS (STRAIN MAASTRICHT).//O12000

F-MAMMA1002880

F-MAMMA1002881//GLIOMA PATHOGENESIS-RELATED PROTEIN (RTVP-1 PROTEIN).//3.3e-22:180:35//HO-MO SAPIENS (HUMAN).//P48060

F-MAMMA1002886//MYOSIN HEAVY CHAIN IB (MYOSIN HEAVY CHAIN IL).//0.00011:148:39//ACANTHAMOE-BA CASTELLANII (AMOEBA).//P19706

F-MAMMA1002887

F-MAMMA1002890//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.030:142:25//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323

F-MAMMA1002892

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F-MAMMA1002895//HYPOTHETICAL PROTEIN UL61.//0.00099:143:35//HUMAN CYTOMEGALOVIRUS (STRAIN AD169),//P16818

F-MAMMA1002908//T-CELL RECEPTOR BETA CHAIN PRECURSOR (ANA 11),//0.12:44:43//ORYCTOLAGUS CUNICULUS (RABBIT),//P06333

F-MAMMA1002909//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00011:28:75//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1002930//BOMBYXIN A-7 PRECURSOR (BBX-A7) (4K-PROTHORACICOTROPIC HORMONE) (4K-PTTH).//0.99:45:46//BOMBYX MORI (SILK MOTH).//P26730

F-MAMMA1002937//ZINC FINGER PROTEIN 42 (MYELOID ZINC FINGER 1) (MZF-1).//6.5e-24:147:34//HOMO SAPIENS (HUMAN),//P28698

F-MAMMA1002938//CERULOPLASMIN PRECURSOR (EC 1.16.3.1) (FERROXIDASE).//4.7e-11:44:68//MUS MUSCULUS (MOUSE).//Q61147

F-MAMMA1002941//PROTEIN Q300.//0.0076:21:61//MUS MUSCULUS (MOUSE).//Q02722

F-MAMMA1002947//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//1.9e-08:152:38//STREP-TOMYCES FRADIAE.//P20186

F-MAMMA1002964

25 F-MAMMA1002970//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//0.0057:55:43//HOMO SAPIENS (HUMAN).// P39189

F-MAMMA1002972//BRAIN-SPECIFIC HOMEOBOX/POU DOMAIN PROTEIN 3A (BRN-3A) (OCT-T1) (HOME-OBOX/POU DOMAIN PROTEIN RDC-1).//0.84:53:41//HOMO SAPIENS (HUMAN).//Q01851

F-MAMMA1002973//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//4.6e-11:54:68//HOMO SAPIENS (HU-MAN).//P39192

F-MAMMA1002982

F-MAMMA1002987//HYPOTHETICAL 11.9 KD PROTEIN IN RPC8-MFA2 INTERGENIC REGION //0.17/47/29// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53906

F-MAMMA1003003//III ALU SUBFAMILY SX WARNING ENTRY IIII//8.6e-09:30:73//HOMO SAPIENS (HU-MAN) //P39195

F-MAMMA1003004 // !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! // 0.0071:41:58 // HOMO SAPIENS (HUMAN). //

F-MAMMA1003007//SPERM\_PROTAMINE\_P1 //0 0076:51:37//TACHYGLOSSUS\_ACULEATUS\_ACULEATUS (AUSTRALIAN ECHIDNA).//P35311

F-MAMMA1003011//HISTONE MACRO-H2A.1.//1.8e-60:175:70//RATTUS NORVEGICUS (RAT).//O02874 F-MAMMA1003013//ACTIN BINDING PROTEIN.//0.097:83:31//SACCHAROMYCES EXIGUUS (YEAST).//

F-MAMMA1003015

F-MAMMA1003019//MYOTUBULARIN.//0.022:56:37//HOMO SAPIENS (HUMAN).//Q13496

45 F-MAMMA1003026//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//0.0014:208:27//ORGYIA PSEUDOTSUGA-TA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//O10341

F-MAMMA1003031//PROBABLE E4 PROTEIN (E1^E4).//0.14:49:32//HUMAN PAPILLOMAVIRUS TYPE 6B.//

F-MAMMA1003035//HYPOTHETICAL 24.4 KD PROTEIN IN LPD 3'REGION (ORF4).//5.1e-12:112:34//ZY-MOMONAS MOBILIS://O66114

F-MAMMA1003039//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/2.4e-07:68:54//HOMO SAPIENS (HUMAN).// P39188

F-MAMMA1003040//!!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!!//2.8e-39:90:57//HOMO SAPIENS (HU-MAN).//P39190

55 F-MAMMA1003044

> F-MAMMA1003047//SPERM HISTONE P2 PRECURSOR (PROTAMINE 2).//0.18:25:44//BOS TAURUS (BO-VINE) //P19782

F-MAMMA1003049//PROBABLE E4 PROTEIN.//0.50:67:29//HUMAN PAPILLOMAVIRUS TYPE 6C.//P20969

- F-MAMMA1003055//WEAK TOXIN CM-2.//0.99:23:30//NAJA HAJE HAJE (EGYPTIAN COBRA).//P01415 F-MAMMA1003056//EXPORTED PROTEIN 7 (FRAGMENT).//1.0:52:32//STREPTOCOCCUS PNEUMONIAE.// P35557
- F-MAMMA1003057//MD6 PROTEIN.//1.5e-85:168:95//MUS MUSCULUS (MOUSE).//Q60584
- F-MAMMA1003066//REGB PROTEIN.//1.0:62:27//PSEUDOMONAS AERUGINOSA.//Q03381
  - F-MAMMA1003089/!!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!!/5.1e-15:44:77//HOMO SAPIENS (HU-MAN)./P39190
    - F-MAMMA1003099//ENDOTHELIAL ACTIN-BINDING PROTEIN (ABP-280) (NONMUSCLE FILAMIN) (FILAMIN 1).//4.8e-20:80:62//HOMO SAPIENS (HUMAN).//P21333
- 10 F-MAMMA1003104//PHOTOSYSTEM I REACTION CENTRE SUBUNIT VIII.//0.98:22:40//SYNECHOCOCCUS ELONGATUS NAEGELI./P25900
  - F-MAMMA1003113//PROCOLLAGEN ALPHA 2(I) CHAIN PRECURSOR (FRAGMENTS).//0.67:35:45//GALLUS GALLUS (CHICKEN).//P02467
- F-MAMMA1003127//MYOSIN I ALPHA (MMI-ALPHA).//5.2e-34:141:56//MUS MUSCULUS (MOUSE).//P46735

  F-MAMMA1003135//HYPOTHETICAL 182.0 KD PROTEIN IN NMD5-HOM6 INTERGENIC REGION.//3.6e-05:91:
  - 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47170
    - F-MAMMA1003146//MALE SPECIFIC SPERM PROTEIN MST87F.//1.0:33:36//DROSOPHILA MELANOGASTER (FRUIT FLY).//P08175
- F-MAMMA1003150//HYPOTHETICAL 84.3 KD PROTEIN ZK945.10 IN CHROMOSOME II.//4.4e-10:254:30// CAENORHABDITIS ELEGANS.//Q09625
  - F-MAMMA1003166//BRAIN PROTEIN H5.//4.0e-42:182:48//HOMO SAPIENS (HUMAN).//O43236
- F-NT2RM1000001//HYPOTHETICAL 8.7 KD PROTEIN IN RPL22-RPL23 INTERGENIC REGION (ORF70).//0.15: 38:34//ASTASIA LONGA (EUGLENOPHYCEAN ALGA).//P34779
- 25 F-NT2RM1000018
  - F-NT2RM1000032//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.51:17:41//CYPRINUS CARPIO (COM-MON CARP).//P24948
  - F-NT2RM1000035//3-HYDROXY-3-METHYLGLUTARYL-COENZYME A REDUCTASE (EC 1.1.1.34) (HMG-COA REDUCTASE).//0.00011:114:27//BLATTELLA GERMANICA (GERMAN COCKROACH).//P54960
- 30 F-NT2RM1000037/I/METALLOTHIONEI/N-II (MT-II)//0.025.19.47/I/SCYLLA SERRATA (MUD CRAB)///P02806 F-NT2RM1000039//VITELLINE MEMBRANE VM34CA PROTEIN PRECURSOR //0.00083:84:33//DROSOPHILA MELANCGASTER (FRUIT FLY)//006521
  - F-NT2RM1000055//HISTIDINE-RICH GLYCOPROTEIN PRECURSOR.//1.1e-07:34:55//PLASMODIUM LOPHU-RAE.//P04929
- 35 F-NT2RM1000059//MYOCYTE-SPECIFIC ENHANCER FACTOR 2B (SERUM RESPONSE FACTOR-LIKE PRO-TEIN 2) (XMEF2) (RSRFR2).//0.18:83:36//HOMO SAPIENS (HUMAN).//Q02080
  - F-NT2RM1000062//PROLINE-RICH PEPTIDE P-B.//0.54:34:44//HOMO SAPIENS (HUMAN).//P02814
  - F-NT2RM1000080//HYPOTHETICAL 35.7 KD PROTEIN SLR1128.//2.1e-20:119:40//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//P72655
- 40 F-NT2RM1000086//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.20:56:35//HOMO SAPIENS (HUMAN).//P10162
  - F-NT2RM1000092//COLLAGEN-LIKE PROTEIN.//0.0017:44:45//HERPESVIRUS SAIMIRI (SUBGROUP C / STRAIN 488).//P22576
- F-NT2RM1000118//CALCINEURIN B SUBUNIT (PROTEIN PHOSPHATASE 2B REGULATORY SUBUNIT) (CALCINEURIN REGULATORY SUBUNIT).//5.7e-07:109:28//NEUROSPORA CRASSA.//P87072
- F-NT2RM1000119//TRANSCRIPTIONAL REGULATOR IE63 (VMW63) (ICP27).//0.0050:135:32//HERPES SIM-PLEX VIRUS (TYPE 2 / STRAIN HG52).//P28276
  - F-NT2RM1000127//EXTENSIN PRECURSOR (PROLINE-RICH GLYCOPROTEIN).//0.032:68:32//SORGHUM VULGARE (SORGHUM).//P24152
- 50 F-NT2RM1000131//METALLOTHIONEIN-III (MT-III) (GROWTH INHIBITORY FACTOR) (GIF).//0.82:33:39//BOS TAURUS (BOVINE).//P37359
  - F-NT2RM1000132//NADH-UBIQUINONE OXIDOREDUCTASE 13 KD-A SUBUNIT PRECURSOR (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-13KD-A) (CI-13KD-A).//2.7e-59:124:91//HOMO SAPIENS (HUMAN).//075380
- F-NT2RM1000153//CYTOSOLIC PURINE 5'-NUCLEOTIDASE (EC 3.1.3.5).//2.5e-08:148:29//HOMO SAPIENS (HUMAN).//P49902
  - F-NT2RM1000186//CALCINEURIN B SUBUNIT (PROTEIN PHOSPHATASE 2B REGULATORY SUBUNIT) (CAL-CINEURIN REGULATORY SUBUNIT).//1.9e-07:109:27//NEUROSPORA CRASSA.//P87072
    - F-NT2RM1000187//PUTATIVE PRE-MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE

- SPAC10F6.02C. //1.0e-12:94:46//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O42643
- F-NT2RM1000199//CUTICLE COLLAGEN 12 PRECURSOR.//0.46:130:33//CAENORHABDITIS ELEGANS.//
- F-NT2RM1000242//PUTATIVE ATP SYNTHASE J CHAIN, MITOCHONDRIAL (EC 3.6.1.34).//0.85:38:36// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//013931
- F-NT2RM1000244//HYPOTHETICAL 131.5 KD PROTÉIN C02F12.7 IN CHROMOSOME X.//0.0055:98:36// CAENORHABDITIS ELEGANS.//O11102
  - F-NT2RM1000252//TRICHOHYALIN.//2.9e-06:88:36//OVIS ARIES (SHEEP).//P22793
- F-NT2RM/100256//GLUCOSAMINE-FRUCTOSE-6-PHOSPHATE AMINOTRANSFERASE [ISOMERIZING]
  10 (EC.2.6.1.16) (HEXOSEPHOSPHATE AMINOTRANSFERASE) (D-FRUCTOSE-6-PHOSPHATE AMIDOTRANSFERASE) (GFATI/Z 9s-54:153:67/I/MUS MUSCULUS (MOUSE)//P47856
  - F-NT2RM1000257//MAGO NASHI PROTEIN://5.9e-64:136:89//DROSOPHILA MELANOGASTER (FRUIT FLY):// P49028

### F-NT2RM1000260

- 15 F-NT2RM1000271//GALACTOKINASE (EC 2.7.1.6).//0.99:41:39//BACILLUS SUBTILIS.//P39574
  - F-NT2RM1000272/HYPOTHETICAL 55.5 KD PROTEIN ZK1128.2 IN CHROMOSOME III.//8.8e-25:131:45// CAENORHABDITIS ELEGANS.//Q09357
    - F-NT2RM1000280//VACUOLAR ATP SYNTHASE SUBUNIT D (EC 3.6.1.34) (V-ATPASE D SUBUNIT) (V- AT-PASE 28 KD ACCESSORY PROTEIN).//2.5e-63:121:94//BOS TAURUS (BOVINE).//P39942
- F-NT2RM1000300//TREACLE PROTEIN (TREACHER COLLINS SYNDROME PROTEIN).//0.51:145:26//HOMO SAPIENS (HUMAN).//Q13428
  - F-NT2RM1000314
  - F-NT2RM1000318//50S RIBOSOMAL PROTEIN L23.//0.83:28:35//AQUIFEX AEOLICUS.//O66433
- F-NT2RM1000341

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- 25 F-NT2RM1000354//HYPOTHETICAL 5.8 KD PROTEIN IN PUHA 5'REGION (ORF55).//0.95:43:37//RHODO-BACTER CAPSULATUS (RHODOPSEUDOMONAS CAPSULATA).//P26159
  - F-NT2RM1000355//SPERM-SPECIFIC PROTEIN PHI-1.//0.0016;73:43//MYTILUS EDULIS (BLUE MUSSEL).// Q04621
  - F-NT2RM1000365//HYPOTHETICAL PROTEIN KIAA0140.//3.5e-10:83:49//HOMO SAPIENS (HUMAN).// Q14153
- U14153
  F-NTZRM1000377//DUAL SPECIFICITY PROTEIN PHOSPHATASE 9 (EC 3.1.3.48) (EC 3.1.3.16) (MITOGEN-ACTIVATED PROTEIN KINASE PHOSPHATASE 4) (MAP KINASE PHOSPHATASE 4) (MKP-4)//4.9e-18:113:38//HOMO SAPIENS (HUMAN)//099956
  - F-NT2RM1000388/HYPOTHETICAL 27.7 KD PROTEIN IN CPT1-SPC98 INTERGENIC REGION.//0.00023:67: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53915
    - F-NT2RM1003394/iHISTONE H3.3 (H3.B) (H3.3Q) /i4.7e-52:71:91/iHOMO SAPIENS (HUMAN), MUS MUSCU-LUS (MOUSE), RATTUS NORYEGICUS (RAT), ORYCTOLAGUS CUNICULUS (RABBIT), GALLUS GALLUS (CHICKEN), SPISULA SOLIDISSIMA (ATLANTIC SURF-CLAM), DROSOPHILA MELANOGASTER (FRUIT FLY), AND DROSOPHILA HYDEI (FRUIT FLY)/iP06351
- 40 F-NT2RM1000399//ENDOTHELIN-2 PRECURSOR (ET-2) (FRAGMENT).//0.92:24:45//CANIS FAMILIARIS (DOG).//P12064
  - F-NT2RM1000421//CUTICLE COLLAGEN 2C (FRAGMENT).//0.12:93:33//HAEMONCHUS CONTORTUS.//
  - F-NT2RM1000430//PISTIL-SPECIFIC EXTENSIN-LIKE PROTEIN PRECURSOR (PELP).//0.13:86:31//NICO-TIANA TABACUM (COMMON TOBACCO).//Q03211
  - F-NT2RM1000499//HYPOTHETICAL PROTEIN KIAA0041 (FRAGMENT).//2.9e-17:75:49//HOMO SAPIENS (HUMAN).//Q15057
    - F-NT2RM1000539//HYPOTHETICAL 10.4 KD PROTEIN IN FTR1-SPT15 INTERGENIC REGION.//2.9e-16:82: 51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40089
  - F-NT2RM1000553//GLYCOLIPID TRANSFER PROTEIN (GLTP).//6.4e-06:103:33//SUS SCROFA (PIG).// P17403
    - F-NT2RM1000555//UNR PROTEIN.//8.7e-77:105:95//RATTUS NORVEGICUS (RAT).//P18395
    - F-NT2RM1000563//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.15:20:50//HO-MO SAPIENS (HUMAN).//P30808
- 55 F-NT2RM1000623//CLARA CELL PHOSPHOLIPID-BINDING PROTEIN PRECURSOR (CCPBP) (CLARA CELLS 10 KD SECRETORY PROTEIN) (CC10).//0.17:70:34//HOMO SAPIENS (HUMAN).//P11684
  - F-NT2RM1000648//GLYCOSYLTRANSFERASE ALG2 (EC 2.4.1.-).//2.0e-22:133:42//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST).//P43636

- F-NT2RM1000661//METALLOTHIONEIN-III (MT-III) (GROWTH INHIBITORY FACTOR) (GIF) (GIFB).//0.0060:24: 33//HOMO SAPIENS (HUMAN).//P25713
- F-NT2RM1000666//COLD SHOCK PROTEIN SCOF.//9.1e-07:67:41//STREPTOMYCES COELICOLOR.//
  P48859
- F-NT2RM1000669//CHLOROPLAST 50S RIBOSOMAL PROTEIN L31.//0.071:69:31//PORPHYRA PURPU-REA.//P51290
  - F-NT2RM1000672//SIGNAL RECOGNITION PARTICLE SEC65 SUBUNIT (FRAGMENT).//0.27:42:42//KLUY-VEROMYCES LACTIS (YEAST).//013475
- F-NT2RM1000691//RETINOBLÁSTOMA BINDING PROTEIN 2 (RBBP-2).//4.3e-42:241:42//HOMO SAPIENS (HUMAN).//P29375
  - F-NT2RM1000699//N2,N2-DIMETHYLGUANOSINE TRNA METHYLTRANSFERASE PRECURSOR (EC 2.1.1.32),//0.94:48:37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P15565
  - F-NT2RM1000702//GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT 1.//0.0013:139:25//DRO-SOPHILA MELANOGASTER (FRUIT FLY).//P26308
- 15 F-NT2RM1000725//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//1.0:15:60//HOMO SAPIENS (HUMAN).// P02811
  - F-NT2RM1000741//STATHMIN (CLONE XO20) (FRAGMENT).//1.0:53:32//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//Q09005
  - F-NT2RM1000742//HYPOTHETICAL 24.1 KD PROTEIN IN DHFR 3'REGION (ORF2).//1.0:54:42//HERPESVI-RUS SAIMIRI (STRAIN 484-77).//P25049
  - F-NT2RM1000746//HYPOTHETICAL 16.8 KD PROTEIN C29E6.04 IN CHROMOSOME I.//0.11:87:21// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09858
    - F-NT2RM1000770//DXS6673E PROTEIN.//2.0e-38:190:48//HOMO SAPIENS (HUMAN).//Q14202
- F-NT2RM1000772//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//4.3e-12:141:30//PODOSPORA AN-SERINA.//Q00808
  - F-NT2RM1000780//MALE SPECIFIC SPERM PROTEIN MST87F.//0.98:34:38//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P08175 F-NT2RM1000781
  - F-NT2RM1000800//24.1 KD PROTEIN IN VMA12-APN1 INTERGENIC REGION.//7.9e-11:135:34//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P28707
  - F-NT2RM1008802/IALPHA-AMYLASE INHIBITOR PAIM I (PIG PANCREATIC ALPHA-AMYLASE INHIBITOR OF MICROBES 1).//0.43:62:35//STREPTOMYCES OLIVACEOVIRIDIS (STREPTOMYCES OCRCHORUSII).// pnoo21

### F-NT2RM1000811

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- 35 F-NT2RM1000826//UNR PROTEIN.//1.1e-110:144:83//RATTUS NORVEGICUS (RAT).//P18395
  - F-NT2RM1000829//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:38:34//DROSOPHILA SIMULANS (FRUIT FLY).//P50270
  - F-NT2RM1000833//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//1.4e-62:145:841/CANIS FA-MILIARIS (DOG).//P38377
- 40 F-NT2RM1000850//TESTIS-SPECIFIC PROTEIN KINASE 1 (EC 2.7.1.-).//6.1e-08:136:33//RATTUS NORVEGI-CUS (RAT).//Q63572
  - F-NT2RM1000852//ATP-DEPENDENT RNA HELICASE ROK1.//1.6e-34:212:43//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P45818
  - F-NT2RM1000857//HISTONE H1.M6.1.//0.76:31:48//TRYPANOSOMA CRUZI.//P40273
- 45 F-NT2RM1000867//MICROSOMAL SIGNAL PEPTIDASE 10.8 KD SUBUNIT (EC 3.4.-.)//0.0082:76:25//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P46965
  - F-NT2RM1000874//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.38:12:58//HO-MO SAPIENS (HUMAN).//P30808
  - F-NT2RM1000882//CYTOCHROME B5.//9.0e-13:92:38//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40312
  - F-NT2RM1000883//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.79:22:59//HO-MO SAPIENS (HUMAN).//P30808
    - F-NT2RM1000885//HYPOTHETICAL 5.8 KD PROTEIN.//0.76:18:38//CLOVER YELLOW MOSAIC VIRUS (CYMV).//P16485
- 59 F.NTZRM1000894/IONA-DIRECTED RNA POLYMERASE 1135 KD POLYPEPTIDE (EC 2.7.7.6) (RNA POLYMERASE I SUBUNIT 2) (RPA135) (RNA POLYMERASE I 127 KD SUBUNIT).//6.2e-70:153:88//RATTUS NORVEGICUS (RAT).//054888
  - F-NT2RM1000898//ACTIN, CYTOPLASMIC (ACTIN, MICRONUCLEAR).//4.3e-12:159:28//OXYTRICHA FAL-

### LAX //P02583

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F-NT2RM1000905//GLUTATHIONE S-TRANSFERASE 1-1 (EC 2.5.1.18) (CLASS-THETA).//0.98:39:35//LUCILIA CUPRINA (GREENBOTTLE FLY) (AUSTRALIAN SHEEP BLOWFLY).//P42860

F-NT2RM1000924//HYPOTHETICAL 39.7 KD PROTEIN C34E10.2 IN CHROMOSOME III.//1.3e-11:169:28// CAENORHARDITIS ELEGANS //P46577

F-NT2RM1000927//CUTICLE COLLAGEN 1.//0.00048:141:31//CAENORHABDITIS ELEGANS.//P08124

F-NT2RM1000962//HYPOTHETICAL 35.8 KD PROTEIN C4F8.04 IN CHROMOSOME I.//7.1e-13:169:31// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O14180

F-NT2RM1000978/HYPOTHETICAL 20.2 KD PROTEIN IN MNN4-PTK1 INTERGENIC REGION.//0.61:82:34// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36045

F-NT2RM1001003//ALPHA-2 CATENIN (ALPHA N-CATENIN) (NEURAL ALPHA-CATENIN).//1.6e-21:211:31// GALLUS GALLUS (CHICKEN).//P30997

F-NT2RM1001008//HYPOTHETICAL 72.5 KD PROTEIN C2F7.10 IN CHROMOSOME I.//3.2e-15:119:36// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09701

15 F-NT2RM1001043/ENDOTHELIN-1 (ET-1) (FRAGMENT),//0.78:32:34//MACACA FASCICULARIS (CRAB EAT-ING MACAQUE) (CYNOMOLGUS MONKEY),//Q28469

F-NT2RM1001044

F-NT2RM1001059//LORICRIN.//8.6e-08:108:39//HOMO SAPIENS (HUMAN).//P23490

F-NT2RM1001066//METALLOTHIONEIN-LIKE PROTEIN TYPE 2.//0.99:24:50//LYCOPERSICON ESCULEN-TUM (TOMATO).//Q43513

F-NT2RM1001072/I1-PHOSPHATIDYLINOSITOL-4,5-BISPHOSPHATE PHOSPHODIESTERASE GAMMA 1 (EC 3.1.4.11) (PLC-GAMMA-1) (PHOSPHOLIPASE C-GAMMA-1) (PLC-II) (PLC-148).//4.7e-15:148:33/HOMO SAPIENS (HUMAN).//P19174

F-NT2RM1001074//HYPOTHETICAL PROTEIN F-215.//8.6e-05:126:30//HUMAN ADENOVIRUS TYPE 2.// P03291

F-NT2RM1001082//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//6.5e-19:75:54//HOMO SAPIENS (HUMAN).//

F-NT2RM1001085/MALE SPECIFIC SPERM PROTEIN MST84DB.//0.49:29:41//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643

30 F-NT2RM1001092//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//2.8e-42:200:38//HOMO SA-PIENS (HUMAN).//P51522

F-NT2RM1001102//HYPOTHETICAL 62.8 KD PROTEIN IN TAF145-YOR1 INTERGENIC REGION.//1.7e-18:161: 36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53331

F-NT2RM1001105//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//4.0e-05:157:35//STREP-TOMYCES FRADIAE.//P20186

F-NT2RM1001112//NONHISTONE CHROMOSOMAL PROTEIN HMG-17.//0.18:20:55//BOS TAURUS (BO-VINE)/P/02313
F-NT2RM1001115

F-NT2RM1001139//GLYCINE-RICH CELL WALL STRUCTURAL PROTEIN 1.8 PRECURSOR (GRP 1.8).//2.0e-25:156:46//PHASEOLUS VULGARIS (KIDNEY BEAN) (FRENCH BEAN) //P10496

F-NT2RM2000006//MITOCHONDRIAL RIBOSOMAL PROTEIN \$12.//0.76:45:35//LEISHMANIA TARENTOLAE (SAUROLEISHMANIA TARENTOLAE).//Q34940

F-NT2RM/2000013//DNA-DIRECTED RNA POLYMERASE III 128 KD POLYPEPTIDE (EC 2.7.7.6) (RNA POLYMERASE III SUBUNIT 2).//3.9e-87:238:65//DROSOPHILA MELANOGASTER (FRUIT FLY).//P25167

45 F-NT2RM2000030//TOXINS 1 AND 2.//0.98:21:42//TRIMERESURUS WAGLERI (WAGLER'S PIT VIPER) (TROPIDOLAEMUS WAGLERI).//P24335

F-NT2RM2000032//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00059:53:49//HOMO SAPIENS (HUMAN).// P39188

F-NT2RM2000042//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//1.0:68:26//HOMO SAPIENS (HUMAN).//P22532

F-NT2RM2000092//HYPOTHETICAL 67.5 KD PROTEIN IN PRPS4-STE20 INTERGENIC REGION.//7.0e-11:80: 40//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38748

F-NT2RM2000093//OVARY MATURATING PARSIN (OMP).//1.0:26:38//LOCUSTA MIGRATORIA (MIGRATORY LOCUST).//P80045

55 F-NT2RM/2000101//HYPOTHETICAL 39.3 KD PROTEIN C02B8.6 IN CHROMOSOME X.//3.3e-09:56:35// CAFNORHARDITIS FLEGANS //O11096

F-NT2RM2000124//CAMP-DEPENDENT PROTEIN KINASE, ALPHA-CATALYTIC SUBUNIT (EC 2.7.1.37) (PKA C-ALPHA).//3.1e-35:77:96//MUS MUSCULUS (MOUSE).//P05132

- F-NT2RM2000191//3',5'-CYCLIC-NUCLEOTIDE PHOSPHODIESTERASE REGA (EC 3.1.4.17) (PDEASE REGA),//3.3e-05:181:27//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD),//Q23917
- F-NT2RM2000192//REPLICATION PROTEIN E1 (FRAGMENTS).//0.019:148:25//COTTONTAIL RABBIT (SHOPE) PAPILLOMAVIRUS (STRAIN WASHINGTON B) (CRPV).//P51894
- 5 F-NT2RM2000239//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.00032:111:32//MUS MUSCULUS (MOUSE).//P05143
  - F-nnnnnnnnnn/METALLOTHIONEIN-LIKE PROTEIN TYPE 2.//0.046:59:33//LYCOPERSICON ESCULEN-TUM (TOMATO).//Q43512
- F-NT2RM2000256/I/GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PRO-10 TEIN) (35 KD LECTIN) (CARBOHYDRATE BINDING PROTEIN 35) (CBP 35) (LAMININ-BINDING PROTEIN) (LECTIN L-29) //0.054-46:34/RATTUS NORVECICUS (RAT),/P08699
  - F-NT2RM2000259//TRANS-ACTING TRANSCRIPTIONAL PROTEIN ICP0 (P135 PROTEIN) (IER 2.9/ER2.6).// 0.27:112:33//BOVINE HERPES VIRUS TYPE 1 (STRAIN JURA).//P29128
- F-NT2RM2000260//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//4.7e-22:191:35//MUS MUSCULUS (MOUSE).//P05143
- F-NT2RM2000287/HYPOTHETICAL 11.8 KD PROTEIN C1B3.02C IN CHROMOSOME I.//5.0e-19:83:53// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O13868
  - F-NT2RM2000322//DIAMINOPIMELATE DECARBOXYLASE (EC 4.1.1.20) (DAP DECARBOXYLASE).//0.47: 117:29//HELICOBACTER PYLORI (CAMPYLOBACTER PYLORI).//P56129
- 20 F-NT2RM2000359//SPORE GERMINATION PROTEIN 270-11.//0.12:83:36//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P22698
  - F-NT2RM2000363//BREAKPOINT CLUSTER REGION PROTEIN.//1.3e-16:203:30//HOMO SAPIENS (HU-MAN)//P11274
- F-NT2RM2000368//DEK PROTEIN.//0.00027:100:32//HOMO SAPIENS (HUMAN).//P35659
- 25 F-NT2RM2000371//POLYRIBONUCLEOTIDE NUCLEOTIDYLTRANSFERASE (EC 2.7.7.8) (POLYNUCLE-OTIDE PHOSPHORYLASE) (PNPASE).//6.8e-36:170:47//ESCHERICHIA COLI.//P05055
  - F-NT2RM2000374//NODAL PRECURSOR.//1.1e-32:64:95//MUS MUSCULUS (MOUSE).//P43021
    F-NT2RM2000395//IMMEDIATE-EARLY PROTEIN IE180.//0.31:41:43//PSEUDORABIES VIRUS (STRAIN INDI-
- ANA-FUNKHAUSER / BECKER) (PRV).//P11675

  F-NT2RM2000402//ENDOSOMAL P24A PROTEIN PRECURSOR (70 KD ENDOMEMBRANE PROTEIN) (PHE-ROMONE ALPHA-FACTOR TRANSPORTER) (ACIDIC 24 KD LATE ENDOCYTIC INTERMEDIATE COMPO-
- NENT).//1.2e-30:228:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32802 F-NT2RM2000407/ITRANSMEMBRANE PROTEIN SEX PRECURSOR.//0.032:105:30//HOMO SAPIENS (HU-MAN)./P51805
- 38 F-NT2RM2000420/METALLOTHIONEIN (MT)./i0.88:42:38//PLEURONECTES PLATESSA (PLAICE).//P07216 F-NT2RM2000422//SODIUM- AND CHLORIDE-DEPENDENT TRANSPORTER NTT73.//2.0e-117:237:87//RAT-TUS NORVEGICUS (RATI-)/094864
  - F-NT2RM2000452//HYPOTHETICAL 63.6 KD PROTEIN IN YPT52-GCN3 INTERGENIC REGION.//1.1e-08:157: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36113
- 40 F-NT2RM2000469//70 KD ANTIGEN //0.050:207:23//SHIGELLA FLEXNERI //P18010

- F-NT2RM2000490//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.022:25:44//HOMO SAPIENS (HUMAN).// P02811
- F-NT2RM2000502//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.0037:17:58//DROSOPHILA MELA-NOGASTER (FRUIT FLY) //001645
- 45 F-NT2RM2000504//HYPOTHETICAL 99.0 KD PROTEIN SPBC119.17.//1.7e-22:195:27//SCHIZOSACCHARO-MYCES POMBE (FISSION YEAST) //O42908
  - F-NT2RM2000522//RAS-RELATED PROTEIN RABA (FRAGMENT).//3.6e-05:67:29//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P34141
  - F-NT2RM2000540//HYPOTHETICAL 83.8 KD PROTEIN C27F2.7 IN CHROMOSOME III.//8.4e-33:214:38// CAENORHABDITIS ELEGANS.//Q18262
  - F-NT2RM2000556//HYPOTHETICAL PROTEIN KIAA0288 (HA6116).//1.7e-09:133:36//HOMO SAPIENS (HU-MAN).//P56524
    - F-NT2RM2000566/INTEGRIN ALPHA-6 PRECURSOR (VLA-6) (CD49F).//2.2e-60:244:51//HOMO SAPIENS (HUMAN).//P23229
- 55 F-NT2RM/2000567//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//2.3e-09:192:34//MUS MUSCULUS (MOUSE).//P05143
  - F-NT2RM2000569//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.0e-08:43:72//HOMO SAPIENS (HUMAN).// P39188

- F-NT2RM2000577//ISOLEUCYL-TRNA SYNTHETASE (EC 6.1.1.5) (ISOLEUCINE--TRNA LIGASE) (ILERS).// 9.1e-54:225:45//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//P73505
- F-NT2RM2000581//SPLICEOSOME ASSOCIATED PROTEIN 49 (SAP 49) (SF3B53).//0.079:111:34//HOMO SA-PIENS (HUMAN).//015427
- F-NT2RM2000588//HYPOTHETICAL PROTEIN KIAA0288 (HA6116).//2.3e-09:193:32//HOMO SAPIENS (HU-
  - F-NT2RM2000594//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.18:33:42//HOMO SAPIENS (HUMAN).//
  - P02811

    F-NT2RM2000599//DNA (CYTOSINE-5)-METHYLTRANSFERASE (EC 2.1.1.37) (DNA METHYLTRANS-
- FERASE) (DNA METASE) (MCMT) (M. MMUI).//1.5e-09:68:45//MUS MUSCULUS (MOUSE).//P13864 F-NT2RM2000609//GRANULIN 2.//0.83:42:35//CYPRINUS CARPIO (COMMON CARP).//P81014
  - F-NT2RM2000612//ZINC FINGER PROTEIN GCS1.//7.2e-05:155:29//SACCHAROMYCES CEREVISIAE (BAK-ERS YEAST).//P36197
- F-NT2RM2000623//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//1.8e-09:196:33//SACCHAROMY-15 CES CEREVISIAE (BAKER'S YEAST).//P32323
- F-NT2RM2000624//SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.070:113:27//DROSOPHILA ERECTA
  - F-NT2RM2000635//SPERM PROTAMINE P1.//0.54:47:38//ANTECHINUS STUARTII.//P42129

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- F-NT2RM2000636//OUTER MEMBRANE PROTEIN H.8 PRECURSOR.//0.096:62:35//NEISSERIA GONOR-RHOEAE.//P11910
- F-NT2RM2000639//HYPOTHETICAL PROTEIN MJ0243.//0.99:32:34//METHANOCOCCUS JANNASCHII.//
  O57694
  - F-NT2RM2000649//NEURONAL CALCIUM SENSOR 1 (NCS-1).//0.00049:70:35//RATTUS NORVEGICUS (RAT), AND GALLUS GALLUS (CHICKEN).//P36610
- 25 F-NT2RM2000669//50S RIBOSOMAL PROTEIN L34.//1.0:34:44//BACILLUS SUBTILIS.//P05647
  - F-NT2RM2000691//ACTIN-LIKE PROTEIN 3 (ACTIN-2).//7.0e-116:243:87//HOMO SAPIENS (HUMAN), AND BOS TAURUS (BOVINE).//P32391
  - F-NT2RM2000714//RAS SUPPRESSOR PROTEIN 1 (RSU-1) (RSP-1 PROTEIN) (RSP-1).//3.8e-21:174:35//HO-MO SAPIENS (HUMAN).//Q15404
- F-NT2RM2000718//HYPOTHETICAL 52.9 KD SERINE-RICH PROTEIN C11G7.01 IN CHROMOSOME I.//0.0022: 174:29//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//013695
  - F-NT2RM2000735//ZINC FINGER PROTEIN 43 (ZINC PROTEIN HTF6).//1.6e-102:246:74//HOMO SAPIENS (HUMAN).//P28160
  - F-NT2RM2000740//HYPOTHETICAL 131.1 KD HELICASE IN ALG7-ENP1 INTERGENIC REGION.//8.5e-51:212: 49//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38144
  - F-NT2RM2000795//!!!! ALU SUBFAMILY SB WARNING ENTRY!!!!//9.0e-41:125:53//HOMO SAPIENS (HU-MAN).//P39189
  - F-NT2RM2000821//COATOMER BETA SUBUNIT (BETA-COAT PROTEIN) (BETA-COP).//1.1e-128:291:89//
    RATTUS NORVEGICUS (RAT).//P23514
- 40 F-NT2RM2000837//CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT KINASE INHIBITOR P57) (P57KIP2).//3.9e-05:113:36//HOMO SAPIENS (HUMAN).//P49918
  - F-NT2RM2000951//HYPOTHETICAL 60.3 KD PROTEIN R08D7.7 IN CHROMOSOME III.//2.5e.49:273:39// CAENORHABDITIS ELEGANS.//P30646
- F-NT2RM2000952//NEUROFILAMENT TRIPLET H PROTEIN (200 KD NEUROFILAMENT PROTEIN) (NF-H)
  (FRAGMENT).//0.037:234:23//RATTUS NORVEGICUS (RAT).//P16884
- F-NT2RM2000984//HYPOTHETICAL 54.7 KD PROTEIN F37A4.1 IN CHROMOSOME III.//6.3e-44:216:43//
  CAENORHABDITIS ELEGANS.//P41879
  - F-NT2RM2001004//SYNAPSINS IA AND IB.//0.15:178:32//RATTUS NORVEGICUS (RAT).//P09951
  - F-NT2RM2001035//CCR4-ASSOCIATED FACTOR 1 (CAF1).//1.4e-87:188:90//MUS MUSCULUS (MOUSE).// Q60809
- F-NT2RM2001065//ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6).//0.53:122:31//TRYPANOSOMA BRU-CEI BRUCEI.//P24499
  - F-NT2RM2001100//HYPOTHETICAL 39.7 KD PROTEIN C34E10.2 IN CHROMOSOME III.//3.4e-13:171:30// CAENORHABDITIS ELEGANS.//P46577
- 55 F-NT2RM2001105//SPORE COAT PROTEIN SP96.//7.8e-06:141:34//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P14328
  - F-NT2RM2001131//PROBABLE EUKARYOTIC INITIATION FACTOR C17C9.03.//2.3e-18:249:31//SCHIZOSAC-CHAROMYCES POMBE (FISSION YEAST).//Q10475

- F-NT2RM2001141//HYPOTHETICAL 115.4 KD PROTEIN ZK757.3 IN CHROMOSOME III.//0.050:134:26// CAENORHABDITIS ELEGANS.//P34681
- F-NT2RM2001152
- F-NT2RM2001177//COLLAGEN ALPHA 1(XIV) CHAIN PRECURSOR (UNDULIN).//0.86:42:40//GALLUS GAL-LUS (CHICKEN) //P32018
- F-NT2RM2001194//SMOOTHELIN.//4.7e-05:77:32//HOMO SAPIENS (HUMAN).//P53814
  - F-NT2RM2001196//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//1.7e-18:218:35//MUS MUSCULUS (MOUSE).//P05143
- F-NT2RM2001201//CYSTEINE STRING PROTEIN (CCCS1).//0.041:22:59//TORPEDO CALIFORNICA (PACIFIC ELECTRIC RAY).//P56101
  - F-NT2RM2001221/KALIRIN (PAM COOH-TERMINAL INTERACTOR PROTEIN 10) (P-CIP10).//1.3e-13:183:32// RATTUS NORVEGICUS (RAT).//P97924
  - F-NT2RM2001238//GLUTAMINASE, KIDNEY ISOFORM PRECURSOR (EC 3.5.1.2) (GLS) (L-GLUTAMINE AMI-DOHYDROLASE).//6.5e-121:218:98//RATTUS NORVEGICUS (RAT).//P13264
- 15 F-NT2RM2001243//HYPOTHETICAL 200.0 KD PROTEIN IN GZF3-IME2 INTERGENIC REGION.//0.00019:177: 27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P42945
  - F-NT2RM2001247//LEGUMIN B (FRAGMENT).//0.22:54:35//PISUM SATIVUM (GARDEN PEA).//P14594
  - F-NT2RM2001256//PROTEIN TSG24 (MEIOTIC CHECK POINT REGULATOR).//1.8e-109:207:98//MUS MUS-CULUS (MOUSE).//P53995
- 20 F-NT2RM2001291//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.016:22:40//HOMO SAPIENS (HUMAN).//P22531
  - F-NTZRMZ001308//REF(2)P PROTEIN./0.61:51:33//DROSOPHILA MELANOGASTER (FRUIT FLY).//P14199 F-NTZRMZ001312//IIII ALU SUBFAMILY SX WARNING ENTRY !III//7.2e-11:33:72//HOMO SAPIENS (HUMAN).// P39196
- 25 F-NT2RM2001319

- F-NT2RM2001324//ZYXIN.//5.1e-22:91:38//GALLUS GALLUS (CHICKEN).//Q04584
- F-NT2RM2001345//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//7.4e-10:159:27//PODOSPORA AN-SERINA.//Q00808
- F-NT2RM2001360//ACCESSORY GLAND PEPTIDE PRECURSOR (PARAGONIAL PEPTIDE B).//1.0:27:48//

  DROSOPHILA MELANOGASTER (FRUIT FLY).//P05623
- F-NT2RM2001370//NAPE PROTEIN.//0.98:44:31//PARACOCCUS DENITRIFICANS (SUBSP. THIOSPHAERA PANTOTROPHA) //056348
  - F-NT2RM2001393//VITELLOGENIN PRECURSOR (VTG) [CONTAINS: LIPOVITELLIN LV-1N; LIPOVITELLIN LV-1C; LIPOVITELLIN LV-2].//0.0024:163:31//ICHTHYOMYZON UNICUSPUS (SILVER LAMPREY).//Q91062
- 35 F-NT2RM2001420 F-NT2RM2001424//HETEROGENOUS NUCLEAR RIBONUCLEOPROTEIN U (HNRNP U).//2.4e-41:140:59//
  - HOMO SAPIENS (HUMAN) //000339
    F-NTZRM2001499//HIGH-AFFINITY CATIONIC AMINO ACID TRANSPORTER-1 (CAT-1) (CAT-1) (SYSTEM Y+BASIC AMINO ACID TRANSPORTER) (ECOTROPIC RETROVIRAL LEUKEMIA RECEPTOR HOMOLOGI/ERR)
- 40 (ECOTROPIC RETROVIRUS RECEPTOR HOMOLOG)//3.7e-71:201:68/HOMO SAPIENS (HUMAN).//P30825 F-NT2RM2001504//CUTICLE COLLAGEN 2.//0.028.41:39//CAENORHABDITIS ELEGANS.//P17656
  - F-NT2RM2001524/HYPOTHETICAL 61.3 KD PROTEIN F25B5.5 IN CHROMOSOME III.//6.7e-47:190:42// CAENORHABDITIS ELEGANS.//Q09316
- F-NT2RM2001544//TELOMERE-BINDING PROTEIN 51 KD SUBUNIT.//0.0027:136:33//EUPLOTES
  5 CRASSUS //O06184
- F-NT2RM2001547//HYPOTHETICAL 48.6 KD PROTEIN IN BET1-PAN1 INTERGENIC REGION.//8.5e-18:91:50// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40564
  - F-NT2RM2001575/52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS-A)) (RO(SS-A)).//3.9e-35:212:41//HOMO SAPIENS (HUMAN).//P19474
  - F-NT2RM2001582//RESA PROTEIN.//0.0033:72:27//BACILLUS SUBTILIS.//P35160
  - F-NT2RM2001588//EXTENSIN PRECURSOR (PROLINE-RICH GLYCOPROTEIN).//1.0e-06:115:32//ZEA MAYS (MAIZE).//P14918
  - F-NT2RM2001592//KERATIN, ULTRA HIGH-SULFUR MATRIX PROTEIN (UHS KERATIN).//0.033:156:23//HO-MO SAPIENS (HUMAN).//P26371
- 55 F-NT2RM2001605//RETINOBLASTOMA BINDING PROTEIN 2 (RBBP-2).//1.1e-116:249:82//HOMO SAPIENS (HUMAN).//P29375
  - F-NT2RM2001613//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//1.2e-97:192:100//RATTUS NORVEGICUS (RAT).//P38378

- F-NT2RM2001632//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.00068:145:28//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P32323
- F-NT2RM2001635//NUCLEAR ENVELOPE PORE MEMBRANE PROTEIN POM 121 (PORE MEMBRANE PROTEIN OF 121 KD) (P145).//1.1e-39:235:47//RATTUS NORVEGICUS (RAT).//P52591
- F-NT2RM2001637//HYPOTHETICAL BHLF1 PROTEIN.//0.075:197:29//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181
  - F-NT2RM2001641//NADH-CYTOCHROME B5 REDUCTASE (EC 1.6.2.2) (B5R).//0.013:29:68//HOMO SAPIENS (HUMAN).//P00387
- F-NT2RM2001648//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//3.2e-65:132:100//CANIS FA-MILIARIS (DOG).//P38377
  - F-NT2RM2001652//PROTEIN TRANSPORT PROTEIN SEC7.//1.6e-32:261:32//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P11075
  - F-NT2RM2001659//CARBOXYPEPTIDASE A INHIBITOR.//0.83:30:46//ASCARIS SUUM (PIG ROUNDWORM)
- 15 F-NT2RM2001664/IKI3 PROTEIN.//1.3e-31:265:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  O06706
  - F-NT2RM2001668//TONB PROTEIN.//0.32:39:41//XANTHOMONAS CAMPESTRIS (PV. CAMPESTRIS).//
  - O34261 F-NT2RM2001670//ZINC FINGER PROTEIN 174.//3.6e-21:172:39//HOMO SAPIENS (HUMAN).//Q15697
- F-NT2RM2001671//HYPOTHETICAL 118.6 KD PROTEIN C29E6.03C IN CHROMOSOME I.//1.6e-10:229:24// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q08857
  FNT2RM2002767(JUNETRODOCHARTE CONTINUED (CALCADED) (PURDED VIA 048424/JUNETRANO
  - F-NT2RM2001675//DIHYDRODIPICOLINATE SYNTHASE (EC 4.2.1.52) (DHDPS).//1.0:184:21//METHANO-COCCUS JANNASCHII.//Q57695
- F-NT2RM2001681//PROTEIN DISULFIDE ISOMERASE PRECURSOR (PDI) (EC 5.3.4.1).//0.0039:199:22//
  DROSOPHILA MELANOGASTER (FRUIT FLY).//P54399
  - F-NT2RM2001688//HYPOTHETICAL 28.1 KD PROTEIN IN SIPU-PBPC INTERGENIC REGION.//2.6e-21:162: 33//BACILLUS SUBTILIS.//P42966
  - F-NT2RM2001695//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//4.9e-41:60:81//HOMO SAPIENS (HUMAN).// P39194
- 30 F-NT2RM2001696/HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION.//9.8e-16:126:38// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41479

- F-NT2RM2001698//PENAEIDIN-3B PRECURSOR (P3-B).//0.36:52:34//PENAEUS VANNAMEI (PENOEID SHRIMP) (EUROPEAN WHITE SHRIMP).//P81059
- F-NT2RM2001699//TRANSCRIPTION INITIATION FACTOR TFIID 30 KD SUBUNIT (TAFII-30) (TAFII30).// 0.0012:79:40//HOMO SAPIENS (HUMAN).//Q12962
- F-NTZRM2001700/IACYL-COA DEHYDROGENASE, VERYL-LONG-CHAIN SPECIFIC (EC 1.3.99.-) (VLCAD) (FNZAMENT)//1.0e-30:140:53//MUS MUSCULLUS (MOUSE).//P50544 F-NTZRM2001706///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.5e-33:95.75//HOMO SAPIENS (HUMAN).//
- P39195
- F-NT2RM/2001716//HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECUR-SOR.//0.010:116:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47179 F-NT2RM/2001718//METHYL-ACCEPTING CHEMOTAXIS PROTEIN TLPB.//0.00029:77:37//BACILLUS SUBTI-
  - LIS.//P39217
  - F-NT2RM2001723//POSTERIOR PITUITARY PEPTIDE.//0.94:26:53//BOS TAURUS (BOVINE).//P01154 F-NT2RM2001727//E7 PROTEIN.//0.91:46:34//HUMAN PAPILLOMAVIRUS TYPE 23.//P50781
- F-NT2RM2001730/PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE K02C4.3 (EC 3.1.2.15) (UBIQUITIN THIOLESTERASE) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE) (DEUBIQUITINATING EN-ZYME)//4.9e-07:39:29/CAENORHABDITIS ELEGANS//0
- F-NT2RM2001743//PROENKEPHALIN A PRECURSOR.//0.75:65:35//CAVIA PORCELLUS (GUINEA PIG).// 50 P47969
  - F-NT2RM2001753//HYPOTHETICAL PROTEIN KIAA0210.//1.5e-14:119:36//HOMO SAPIENS (HUMAN).// Q92609
  - F-NT2RM2001760//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//8.3e-58:119:99//CANIS FA-MILIARIS (DOG).//P38377
- 55 F-NT2RM2001768//HYPOTHETICAL PROTEIN UL25.//0.45:77:32//HUMAN CYTOMEGALOVIRUS (STRAIN AD169) //P16761
  - F-NT2RM2001771//ZINC FINGER PROTEIN 135./I/4.6e-80:224:60//HOMO SAPIENS (HUMAN).//P52742 F-NT2RM2001782//MANNOSE-1-PHOSPHATE GUANYLTRANSFERASE (EC 2.7.7.13) (ATP-MANNOSE-1-

- PHOSPHATE GUANYLYLTRANSFERASE) (NDP-HEXOSE PYROPHOSPHORYLASE).//7.0e-06:61:45//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P41940
- F-NT2RM2001784//HYPOTHETICAL PROTEIN UL61.//0.00070:145:33//HUMAN CYTOMEGALOVIRUS (STRAIN AD169).//P16818
- F-NT2RM2001785//LINOLEOYL-COA DESATURASE (EC 1.14.99.25) (DELTA(6)-DESATURASE).//1.5e-08:127: 32//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//Q08871
  - F-NT2RM2001797//ZINC FINGER PROTEIN 135.//1.6e-73:267:49//HOMO SAPIENS (HUMAN).//P52742 F-NT2RM2001800//HYPOTHETICAL HELICASE MG018/MG017/MG016 HOMOLOG.//3.9e-12:171:33//MYCO-
- PLASMA PNEUMONIAE./IP75093
  for F-NT2RM2001803//IKI3 PROTEIN.//1.6e-38:283:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  OG6706
  - F-NT2RM2001805//COLD SHOCK-LIKE PROTEIN CSPH.//0.51:46:32//SALMONELLA TYPHIMURIUM.//
    O33793
  - F-NT2RM2001813//HYPOTHETICAL 40.4 KD TRP-ASP REPEATS CONTAINING PROTEIN C14B1.4 IN CHRO-
- MOSOME III.//5.0e-05:82:32//CAENORHABDITIS ELEGANS.//Q17963 F-NT2RM2001823//CHROMODOMAIN-HELICASE-DNA-BINDING PROTEIN 2 (CHD-2).//3.6e-49:233:45//HO-
  - MO SAPIENS (HUMAN)//014647
    F-NTZRM2001839//RETICULOCALBIN 1 PRECURSOR.//5 2e-65:222:56//HOMO SAPIENS (HUMAN).//015283
    F-NTZRM2001840///III ALU SUBFAMILY SQ WARNING ENTRY!/!!///9 6e-33:102:68//HOMO SAPIENS (HU-
  - MAN).//P39194 F-NT2RM2001855//BASP1 PROTEIN.//0.054;120;30//HOMO SAPIENS (HUMAN).//P80723

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- F-NT2RM2001867//HYPOTHETICAL 56.6 KD PROTEIN IN URE2-SSU72 INTERGENIC REGION.//4.1e-19:88: 36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53867
- F-NT2RM2001879//HYPOTHETICAL 47.3 KD PROTEIN C22G7.07C IN CHROMOSOME I.//5.9e-15:76:38// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//009800
  - F-NT2RM2001886//HYPOTHETICAL 126.9 KD PROTEIN C22G7.04 IN CHROMOSOME I.//1.4e-41:249:38//
    SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//1009798
    - F-NT2RM2001896//HYPOTHETICAL 83.2 KD PROTEIN IN KAR4-PBN1 INTERGENIC REGION.//2.1e-59:197: 56//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P25582
- 30 F-NT2RM2001903//HYPOTHETICAL PROTEIN MJ0263.//0.070:132:31//METHANOCOCCUS JANNASCHII.// O06917
  - F-NT2RM2001930//THROMBOSPONDIN 2 PRECURSOR.//7.1e-05:53:47//MUS MUSCULUS (MOUSE).//
    - F-NT2RM2001935//PUTATIVE CUTICLE COLLAGEN F55C10.3.//0.00046:116:35//CAENORHABDITIS ELE-GANS.//Q21184
      - F-NT2RM2001936//32.3 KD PROTEIN IN CWP1-MBR1 INTERGENIC REGION.//4.5e-27:216:34//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P28320
      - F-NT2RM2001950//HIRUDIN HV1 (BUFRUDIN).//0.59:43:34//HIRUDINARIA MANILLENSIS (BUFFALO LEECH).///981492
- 40 F-NT2RM2001982//GUANINE NUCLEOTIDE-BINDING PROTEIN G(I)/G(S)/G(O) GAMMA-8 SUBUNIT (G GAM-MA-C).//0.72:35:42//BOS TAURUS (BOVINE).//P50154
  - F-NT2RM2001983//PROLINE-RICH PEPTIDE P-B.//0.00035:23:52//HOMO SAPIENS (HUMAN).//P02814 F-NT2RM2001989//NUCLEOLAR PROTEIN NOP4 (NUCLEOLAR PROTEIN NOP77\.//8.6e-24:197:37//SAC-
- F-N12KM2001989/MUCLEOLAN PROTEIN NOP4 (NUCLEOLAN PROTEIN NOP77)///8.68-24:197:37//SAC CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P37838 45 F-NTZRM2001997
- F-NT2RM2001998//IMMEDIATE-EARLY PROTEIN IE180.//0.076:92:27//PSEUDORABIES VIRUS (STRAIN IN-DIANA-FUNKHAUSER / BECKER) (PRV).//P11675
  - F-NT2RM2002004//SLF1 PROTEIN.//3.5e-06:235:27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// Q12034
- F-NT2RM2002014//HYPOTHETICAL PROTEIN HI0568.//2.1e-17:235:29//HAEMOPHILUS INFLUENZAE.// P71353
  - F-NT2RM2002030/IGLUCOSAMINE-FRUCTOSE-8-PHOSPHATE AMINOTRANSFERASE [ISOMERIZING] (EC 2.6.1.16) (HEXOSEPHOSPHATE AMINOTRANSFERASE) (D-FRUCTOSE-8- PHOSPHATE AMIDOTRANSFERASE) (G-FATI//9.5-9-105.271-78/MUS MUSCULUS (MOUSE)//P47856
- 55 F-NT2RM2002049/SMALL PROLINE-RICH PROTEIN 2-1/1/0.699-41-41/IHOMO SAPIENS (HUMAN)/IP35326 F-NT2RM2002055//ACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS13./I/0.012/217.24//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST)./I/007878
  - F-NT2RM2002088//PUTATIVE HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN X (HNRNP X) (CBP).//

1.1e-09:65:53//MUS MUSCULUS (MOUSE) //Q61990

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- F-NT2RM2002091//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.072:74: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214
- F-NT2RM2002100//ATP-DEPENDENT RNA HELICASE ROK1.//4.5e-50:289:41//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P45818
- F-NT2RM2002109//NT-3 GROWTH FACTOR RECEPTOR PRECURSOR (EC 2.7.1.112) (TRKC TYROSINE KINASE) (GP145-TRKC) (TRK-C).//1.4e-14:203:32//RATTUS NORVEGICUS (RAT).//Q03351
- F-NT2RM2002128//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//0.0025:139:31// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437
- 10 F-NT2RM2002142//GASTRULATION SPECIFIC PROTEIN G12.//9.2e-20:42:73//BRACHYDANIO RERIO (ZE-BRAFISH) (ZEBRA DANIO).//P47805
  - F-NT2RM2002145//GLUTENIN, HIGH MOLECULAR WEIGHT SUBUNIT 12 PRECURSOR.//0.0085;200:26// TRITICUM AESTIVUM (WHEAT).//P08488
  - F-NT2RM2002178//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//5.8e-05:56:39//BOS TAURUS (BO-VINE).//P25508
  - F-NT2RM2002580//CCAAT-BINDING TRANSCRIPTION FACTOR SUBUNIT A (CBF-A) (NF-Y PROTEIN CHAIN B) (NF-YB) (CAAT-BOX DNA BINDING PROTEIN SUBUNIT B) //2-96-14-96-37//PETROMYZON MARINUS (SEA LAMPREY) ///25210
- F-NT2RM4000024//DNA-DIRECTED RNA POLYMERASE III 128 KD POLYPEPTIDE (EC 2.7.7.6) (RNA POLYMERASE III SUBUNIT 2)//8.69-95:271:67//DROSOPHILA MELANOGASTER (FRUIT FLY)//P25167 F-NT2RM4000027/INTTERFERON-ACTIVATABLE PROTEIN 202 (IFI-202)//0.99-72:31/MUS MUSCULUS
  - (MOUSE).//P15091
    F-NT2RM4000030//LAS1 PROTEIN./1.4e-14:184:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
- P36146
  25 F-NT2RM4000046//COLLAGEN ALPHA 1(III) CHAIN (FRAGMENT).//0.99:120:28//RATTUS NORVEGICUS (RAT).//P13941
- F-NTZRM4000061 F-NTZRM4000085//ATP-DEPENDENT RNA HELICASE A (NUCLEAR DNA HELICASE II) (NDH II) (DEAD BOX
- PROTEIN 9) (MHEL-5).//8.5e-40:263:38//MUS MUSCULUS (MOUSE).//O70133
  30 F-NT2RM4000086//HYPOTHETICAL PROTEIN HI1497.//10.227:37//HAEMOPHILUS INFLUENZAE.//P44221
  - F-NT2RM4000104/ZINC FINGER PROTEIN 134 //1.0e-26:64:56//HOMO SAPIENS (HUMAN)//P52741 F-NT2RM4000139//PREPROTEIN TRANSLOCASE SECE SUBUNIT.//0.99:38:42//THERMOTOGA MARITIMA.// P35874
  - F-NT2RM4000155/THREONYL-TRNA SYNTHETASE, CYTOPLASMIC (EC 6.1.1.3) (THREONINE--TRNA LIGASE) (THRRS).//6.3e-34:181:40//HOMO SAPIENS (HUMAN)//P26639
    - F-NT2RM4000156//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).//
      4.6e-12:142:33//NICOTIANA TABACUM (COMMON TOBACCO).//P13983
    - F-NT2RM4000167//KINESIN-LIKE PROTEIN KIF4.//3.4e-123:269:91//MUS MUSCULUS (MOUSE).//P33174 F-NT2RM4000168/JM PROTEIN, SEROTYPE 2.2 PRECURSOR.//9.7e-10.229:26//STREPTOCOCCUS PYO-GENES.//P60469
- F-NT2RM4000191//P68-LIKE PROTEIN.//2.1e-11:104:40//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P24783
  - F-NT2RM4000197//CUTICLE PROTEIN CP463 (CPCP463).//0.84:29:37//CANCER PAGURUS (ROCK CRAB).//
- 45 F-NT2RM4000199//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//1.8e-06:187:34//HOMO SAPIENS (HUMAN).//P10162
  - F-NT2RM4000200//HYPOTHETICAL 9.4 KD PROTEIN IN FLAL 3'REGION (ORF3).//0.52:42:40//BACILLUS LI-CHENIFORMIS.//P22754
  - F-NT2RM4000202//COLLAGEN ALPHA 1(VIII) CHAIN PRECURSOR (ENDOTHELIAL COLLAGEN).//0.00044: 168:32//ORYCTOLAGUS CUNICULUS (RABBIT).//P14282
  - F-NT2RM4000210//EXTENSIN PRECURSOR.//0.27:129:27//DAUCUS CAROTA (CARROT).//P06599
    F-NT2RM4000215//MAK16 PROTEIN.//2.0e-65:234:52//SACCHAROMYCES CEREVISIAE (BAKER'S YFAST).//P106872
- F-NT2RM4000229//GAR2 PROTEIN.//0.13:217:26//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//
  P41891
- F-NT2RM4000233/TRANSMEMBRANE PROTEIN SEX PRECURSOR.//0.047:108:30//HOMO SAPIENS (HU-MAN).//P51805
  - F-NT2RM4000244//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.67:59:27//BALAENOPTERA

PHYSALUS (FINBACK WHALE) (COMMON RORQUAL) //P24947

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- F-NT2RM4000251//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.0059:108:35//MUS MUSCULUS (MOUSE).//P05143
- F-NT2RM4000265//IIII ALU SUBFAMILY J WARNING ENTRY IIII//8.1e-38:70:70//HOMO SAPIENS (HUMAN).// P39188
- F-NT2RM4000290//TRANSDUCIN-LIKE ENHANCER PROTEIN 3 (ESG3).//1.6e-115:209:94//HOMO SAPIENS (HUMAN).//O04726
  - F-NT2RM4000324//PRESPORE PROTEIN DP87 PRECURSOR.//0.14:136:30//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//Q04503
- 10 F-NT2RM4000327//HYPOTHETICAL 8.9 KD PROTEIN IN IEO-IE1 INTERGENIC REGION.//0.91:73:28// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41703
  - F-NT2RM4000344//YME1 PROTEIN HOMOLOG (EC 3.4.24.-).//9.4e-78:241:55//CAENORHABDITIS ELE-GANS.//P54813
- F-NT2RM4000349//CYSTEINE STRING PROTEIN (CCCS1).//0.055:22:59//TORPEDO CALIFORNICA (PACIFIC LECTRIC RAY).//P56101
- F-NT2RM4000354//LETHAL(2)DENTICLELESS PROTEIN (DTL83 PROTEIN).//4.6e-26:208:35//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q24371
  - F-NT2RM4000356//COAT PROTEIN.//0.11:105:36//SATELLITE TOBACCO MOSAIC VIRUS (STMV).//P17574 F-NT2RM4000366//IMMEDIATE-EARLY PROTEIN.//1.2e-05:215:24//HERPES VIRUS SAIMIRI (STRAIN 11).//
  - F-NT2RM4000366//IMMEDIATE-EARLY PROTEIN.//1.2e-05:215:24//HERPES VIRUS SAIMIRI (STRAIN 11). Q01042
    - F-NT2RM4000368//HYPOTHETICAL 7.3 KD PROTEIN IN RPBA-GP46 INTERGENIC REGION.//0.54:46:36// BACTERIOPHAGE RB69.//O64300
      - F-NT2RM4000386//RHSC PROTEIN PRECURSOR.//0.0096:162:29//ESCHERICHIA COLI.//P16918
- F-NT2RM4000395//HYPOTHETICAL 52.9 KD PROTEIN IN SAP155-YMR31 INTERGENIC REGION.//4.5e-66: 256:53//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43616
- F-NT2RM4000414//HYPOTHETICAL 6.0 KD PROTEIN IN THI12 5'REGION.//0.13:33:48//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53820
- F-NT2RM4000421//MRNA TRANSPORT REGULATOR MTR10.//5.0e-13:171:29//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//Q99189
- 30 F-NT2RM4000425//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.1e-25:46:80//HOMO SAPIENS (HUMAN).// P39193
  - F-NT2RM4000433//CUTICLE COLLAGEN 3A3./I/2.5e-06:77:38//HAEMONCHUS CONTORTUS.//P16253 F-NT2RM4000457//HYPOTHETICAL 111.9 KD PROTEIN C22H10.03C IN CHROMOSOME I.//4.3e-09:215:22//
- SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10297
  35 F-NT2RM4000471//TRNA SPLICING PROTEIN SPL1.//6.7e-73:163:65//CANDIDA ALBICANS (YEAST).//
  - F-NT2RM4000486//COLLAGEN ALPHA 2(VI) CHAIN PRECURSOR.//0.0012:121:34//GALLUS GALLUS (CHICKEN).//P15988
- F-NT2RM-000496//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RNA POLYMER-49 ASE II SUBUNIT 1)://5.9e-09:175:35//SCH/ZOSACCHAROMYCES POMBE (FISSION YEAST)://P36594
- F-NT2RM4000511//SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.020:122:31//DROSOPHILA SIMU-LANS (FRUIT FLY)//P13729
  - F-NT2RM4000514//ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6),//0.46:68:32//ARTEMIA SANFRAN-CISCANA (BRINE SHRIMP) (ARTEMIA FRANCISCANA),//Q37708
- 45 F-NT2RM4000515//GAR2 PROTEIN.//3.2e-05:198:27//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P41891
- F-NT2RM4000520//HYPOTHETICAL 7.5 KD PROTEIN (ORF 63).//0.011:55:38//SPINACIA OLERACEA (SPINACH).//P08974
- F-NT2RM4000531//ZINC FINGER PROTEIN 169 (FRAGMENT).//3.6e-44:244:42//HOMO SAPIENS (HUMAN).//
- F-NT2RM4000532//PUTATIVE MEMBRANE PROTEIN 53.//1.0:47:34//HERPES VIRUS SAIMIRI (STRAIN 11).//
  - F-NT2RM4000534//HYPOTHETICAL 5.9 KD PROTEIN IN WRBA-PUTA INTERGENIC REGION.//0.75:26:46// ESCHERICHIA COLI.//P56614
- F-NT2RM4000585//GAG POLYPROTEIN [CONTAINS: CORE PROTEIN P16; CORE PROTEIN P26].//0.019:86: 34//HUMAN IMMUNODEFICIENCY VIRUS TYPE 2 (ISOLATE SBLIS Y) (HIV-2).//P12450
  - F-NT2RM4000590//RING CANAL PROTEIN (KELCH PROTEIN).//5.0e-23:224:29//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652

- F-NT2RM4000595//HYPOTHETICAL 54.9 KD PROTEIN C02F5.7 IN CHROMOSOME III.//3.8e-62:226:50// CAENORHARDITIS ELEGANS //P34284
- F-NT2RM4000603//SRC SUBSTRATE CORTACTIN (AMPLAXIN) (EMS1 ONCOGENE).//0.077:132:22//HOMO SAPIENS (HUMAN).//O14247
- F-NT2RM4000611//HYPOTHETICAL 40.4 KD TRP-ASP REPEATS CONTAINING PROTEIN C14B1.4 IN CHRO-MOSOME III.//1.9e-06:82:32//CAENORHABDITIS ELEGANS.//Q17963
  - F-NT2RM4000616//ACETYL-COENZYME A SYNTHETASE (EC 6.2.1.1) (ACETATE--COA LIGASE) (ACYL- AC-TIVATING ENZYME).//5.3e-79:213:62//ESCHERICHIA COLI.//P27550
  - F-NT2RM4000674//HYPOTHETICAL SYMPORTER SLL 1374.//1.3e-11:147:32//SYNECHOCYSTIS SP. (STRAIN
- 10 PCC 6803).//P74168 F-NT2RM4000689
  - F-NT2RM4000698//CHORION CLASS HIGH-CYSTEINE HCA PROTEIN 12 PRECURSOR (HC-A.12).//0.26:45: 33//BOMBYX MORI (SILK MOTH).//P05687
  - F-NT2RM4000700//THIOPHENE AND FURAN OXIDATION PROTEIN THDF.//0.95.165:25//BORRELIA BURG-DORFERI (LYME DISEASE SPIROCHETE).//P53364
- DORFERI (LYME DISEASE SPIROCHETE)./PS3364 F-NT2RM4000712//PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE R10E11.3 (EC 3.1.2.15) (UBIQUITIN THIOLESTERASE) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE) (DEUBIQUITINATING EN-
  - ZYME).//2.2e-82:152:63://CAENORHABDITIS ELEGANS.//P34547
    F-NT2RM4000717i/SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.80:54:40//DROSOPHILA SIMULANS
    (FRIIT FLY // IP)372:0
    - F-NT2RM4000733//OCTAPEPTIDE-REPEAT PROTEIN T2.//1.5e-08:139:28//MUS MUSCULUS (MOUSE).//
    - F-NT2RM4000734//GASTRULA ZINC FINGER PROTEIN XLCGF26.1 (FRAGMENT).//7.2e-20:205:28//XENO-PUS LAEVIS (AFRICAN CLAWED FROG).//P18715
- 25 F-NT2RM4000741//SPERM PROTAMINE P1.//0.89:52:38//ISOODON MACROURUS (SHORT-NOSED BANDI-COOT).//P42136
  - F-NT2RM4000751//ZINC FINGER PROTEIN 26 (ZFP-26) (MKR3 PROTEIN) (FRAGMENT).//5.2e-77:246:52// MUS MUSCULUS (MOUSE).//P10076
  - F-NT2RM4000764//KERATIN, GLYCINE/TYROSINE-RICH OF HAIR://0.062:33:42//OVIS ARIES (SHEEP)://
- F-NT2RM4000778

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- F-NT2RM4000779//SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185).//0.014:53:45//VOLVOX CARTERI.// P21997
- F-NT2RM4000787//BONE MORPHOGENETIC PROTEIN 1 PRECURSOR (EC 3.4.24.-) (BMP-1).//0.00011:73: 39//MUS MUSCULUS (MOUSE).//P98063
- F-NT2RM4000790//SPORE COAT PROTEIN SP96.//0.00083:157:29//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P14328
- F-NT2RM4000795//CHOLINESTERASE PRECURSOR (EC 3.1.1.8) (ACYLCHOLINE ACYLHYDROLASE) (CHOLINE ESTERASE) (IN (BUTYRYLCHOLINE ESTERASE) (PSEUDOCHOLINESTERASE) ///7.4e-41:271:36// HOMO SAPIENS (HUMAN) //P08276
  - F-NT2RM4000796//5-METHYLCYTOSINE-SPECIFIC RESTRICTION ENZYME B (EC 3.1.21.-).//0.28:82:30//ES-CHERICHIA COLL //P15005
  - F-NT2RM4000798//PROTEIN TRANSPORT PROTEIN SEC7.//4.7e-38:165:48//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P11075
- 45 F-NT2RM4000813//METALLOTHIONEIN-IB.//0.0025:25:44//OVIS ARIES (SHEEP).//P09577
  - F-NT2RM4000820
    - F-NT2RM4000833//HYPOTHETICAL PROTEIN MJ1136.//6.5e-42:206:41//METHANOCOCCUS JANNASCHII.// Q58536
    - F-NT2RM4000848//BRAIN-SPECIFIC HOMEOBOX/POU DOMAIN PROTEIN 3A (BRN-3A) (BRN-3.0).//0.00060: 159:33//MUS MUSCULUS (MOUSE).//P17208
    - 199:33//MUS MUSCULUS (MUUSE)//P1/200
      F-NT2RM4000852//SMALL PROLINE-RICH PROTEIN 2B (SPR-2B).//0.0076:13:69//HOMO SAPIENS (HU-MAN)//P35325
      - F-NT2RM4000855//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//0.0060:68:44//HOMO SAPIENS (HUMAN).// P39194
- 55 F-NT2RM4000887//RTS1 PROTEIN (SCS1 PROTEIN).//0.23:153:24//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P38903
  - F-NT2RM4000895/HYPOTHETICAL 53.5 KD PROTEIN IN PHO2-POL3 INTERGENIC REGION.//3.3e-09:80: 46//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43123

- F-NT2RM4000950//HYPOTHETICAL PROTEIN MJ0572.//0.090:68:29//METHANOCOCCUS JANNASCHII.// Q57992
- F-NT2RM4000971/KINESIN LIGHT CHAIN (KLC)//0.79:201:24/ILOLIGO PEALEII (LONGFIN SQUID).//P46825 F-NT2RM4000979//MYOSIN REGULATORY LIGHT CHAIN 2, NONSARCOMERIC (MYOSIN RLC).//1.2e-07:25: 96//HOMO SAPIENS (HUMAN).//P19105
- F-NT2RM4000996/ZINC FINGER PROTEIN 37 (ZFP-37) (MALE GERM CELL SPECIFIC ZINC FINGER PRO-TEIN)//1.4e-56:253:46/MUS MUSCULUS (MOUSE).//P17141 F-NT2RM4001002
- F-NT2RM4001016//GAG POLYPROTEIN [CONTAINS: CORE PROTEIN P15; INNER COAT PROTEIN P12;
  - CORE SHELL PROTEIN P30].//0.25:101:31//FBR MURINE OSTEOSARCOMA VIRUS.//P29175
    - F-NT2RM4001032//CUTICLE COLLAGEN 2.//2.6e-07:130:39//CAENORHABDITIS ELEGANS.//P17656
    - F-NT2RM4001047//MO25 PROTEIN://5.6e-107:252:80//MUS MUSCULUS (MOUSE)://Q06138

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- F-NT2RM4001054//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//9.0e-109:209:94//CANIS FA-MILIARIS (DOG).//P38377
- 15 F-NT2RM4001084//HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN UXUR-IADA INTERGENIC RE-GION.//0.57:95:30//ESCHERICHIA COLI.//P39376
  - F-NT2RM4001092/HYPOTHETICAL 127.4 KD PROTEIN F07F6.4 IN CHROMOSOME III.//2.5e-47:231:47// CAENORHABDITIS ELEGANS.//Q09531
  - F-NT2RM4001116//HYPOTHETICAL 216.3 KD PROTEIN R06F6.8 IN CHROMOSOME II.//1.3e-08:243:23// CAENORHABDITIS ELEGANS.//Q09417
  - F-NT2RM4001140//HOMEOBOX PROTEIN MSH-D.//7.1e-13:103:38//BRACHYDANIO RERIO (ZEBRAFISH) (ZEBRA DANIO).//Q01704
  - F-NT2RM4001151//SYNAPSINS IA AND IB (BRAIN PROTEIN 4.1).//0.26:96:34//HOMO SAPIENS (HUMAN).// P17600
- 25 F-NT2RM4001155//ADRENAL MEDULLA 50 KD PROTEIN.//3.6e-103:201:91//BOS TAURUS (BOVINE).// Q27969
  - F-NTZRM4001160/IGLUTATHIONE S-TRANSFERASE (EC 2.5.1.18) (CLASS-PHI) (FRAGMENTS)//1.0.33:36// BRASSICA OLERACEA (CAULIFLOWER)//P48438 F-NTZRM4001187//PREPROTEIN TRANSLOCASE SECA SUBUNTI/10.44:158.27/MYCOPLASMA GENITALIUM://P47318
- 30 F-NT2RM4001191/LONG NEUROTOXIN 2 (TOXIN C).//0.99:44:43//ASTROTIA STOKESI (STOKES'S SEA SNAKE) (DISTEIRA STOKESI).//P01381
  - F-NT2RM4001200//ZINC FINGER PROTEIN 135.//2.2e-82:245.59//HOMO SAPIENS (HUMAN).//P52742 F-NT2RM4001220//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.028:94: 40//SACCHAROMYCES CEREVISIAE (RAKER'S YEAST)/P58214
- 35 F-NT2RM4001204//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//0.0096:182:34//HOMO SAPIENS (HUMAN).//Q15428
  - F-NT2RM4001217//RING CANAL PROTEIN (KELCH PROTEIN).//2.1e-21:221:29//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- F-NT2RM4001256//CBP3 PROTEIN PRECURSOR.//0.30:55:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P21560
- F-NT2RM4001258//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.00031:132:39//STREP-TOMYCES FRADIAE.//P20186
  - F-NT2RM4001309//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONE CP7) [CONTAINS: BASIC PEPTIDE P-F] (FRAGMENT).//0.048:132:28//HOMO SAPIENS (HUMAN).//P02812
- 45 F-NT2RM4001313//PHOSPHATIDYLINOSITOL 3-KINASE VPS34-LIKE (EC 2.7.1.137) (PI3-KINASE) (PTDINS-3-KINASE) (PI3K).//2.6e-37.124:65//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P54676
  - F-NT2RM4001316//ACYL-COA DEHYDROGENASE, MEDIUM-CHAIN SPECIFIC PRECURSOR (EC 1.3.99.3) (MCAD).//1.7e-10:185:30//RATTUS NORVEGICUS (RAT).//P08503
  - F-NT2RM4001320//PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR (RHO/RAC GEF) (FA-CIOGENITAL DYSPLASIA PROTEIN HOMOLOG).//1.5e-08:197:26//MUS MUSCULUS (MOUSE).//P52734 F-NT2RM4001340//UTR4 PROTEIN (UNKNOWN TRANSCRIPT 4 PROTEIN)//7.7e-14:82:36//SACCHAROMY-
    - F-NT2RM4001344//HYPOTHETICAL GTP-BINDING PROTEIN IN POP2-HOL1 INTERGENIC REGION.//3.3e-16:128:35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53742
- 55 F-NT2RM4001347//HYPOTHETICAL 76.9 KD PROTEIN IN RPM2-TUB1 INTERGENIC REGION.//0.067:111:33// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//QQ4511 E-NT2RM4001371

CES CEREVISIAE (BAKER'S YEAST).//P32626

F-NT2RM4001382//HISTIDINE-RICH GLYCOPROTEIN PRECURSOR.//1.0e-08:82:39//PLASMODIUM LOPHU-

- RAE //P04929
- F-NT2RM4001384
- F-NT2RM4001410//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR. I/2.1e-08:185:31//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323
- F-NT2RM4001411//EARLY NODULIN 20 PRECURSOR (N-20).//5.3e-05:105:38//MEDICAGO TRUNCATULA (BARREL MEDIC),//P93329
  - F-NT2RM4001412//GTPASE-ACTIVATING PROTEIN (GAP) (RAS P21 PROTEIN ACTIVATOR) (P120GAP) (RASGAP).//6.2e-17:109:41//RATTUS NORVEGICUS (RAT).//P50904
  - F-NT2RM4001414//ZINC FINGER PROTEIN 177.//8.3e-06:54:50//HOMO SAPIENS (HUMAN).//Q13360
- 10 F-NT2RM4001437//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//2.1e-24:87:65//HOMO SAPIENS (HUMAN).// P39192
  - F-NT2RM4001444//PROBABLE ISOLEUCYL-TRNA SYNTHETASE (EC 6.1.1.5) (ISOLEUCINE--TRNA LIGASE) (ILERS) (FRAGMENT).//2.6e-45:197:47//CIONA INTESTINALIS.//Q94425
- F-NT2RM4001454//HYPOTHETICAL PROTEIN KIAA0041 (FRAGMENT), I/0, 0060:95;29//HOMO SAPIENS (HU-15 MAN).//Q15057
- F-NT2RM4001455//PROBABLE E5B PROTEIN.//0.41:44:36//HUMAN PAPILLOMAVIRUS TYPE 6B.//P06461
  - F-NT2RM4001483//ZINC FINGER PROTEIN 136.//1.7e-28:85:64//HOMO SAPIENS (HUMAN).//P52737 F-NT2RM4001489//PTB-ASSOCIATED SPLICING FACTOR (PSF).//0.086:111:34//HOMO SAPIENS (HUMAN).//
- 20 F-NT2RM4001519//ACID UREASE ALPHA SUBUNIT (EC 3.5.1.5) (UREA AMIDOHYDROLASE).//0.82:51:47// LACTOBACILLUS FERMENTUM.//P26929
- F-NT2RM4001522//TROPOMYOSIN.//0.030:117:23//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).// 002088

### F-NT2RM4001557

- 25 F-NT2RM4001565//HYPOTHETICAL 44.3 KD PROTEIN C1F7.07C IN CHROMOSOME 1.//0.99:42:40// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09919
  - F-NT2RM4001566//GLUCOAMYLASE S1/S2 PRECURSOR (EC 3,2,1,3) (GLUCAN 1,4-ALPHA- GLUCOSI-DASE) (1,4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//0.054:190:23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P08640
- F-NT2RM4001569//GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT-LIKE PROTEIN (ACTIVATED PROTEIN KINASE C RECEPTOR HOMOLOG).//0.72:64:31//TRYPANOSOMA BRUCEI BRUCEI.//Q94775 E-NIT2RMAN01582
  - F-NT2RM4001592//DNA REPAIR PROTEIN RAD9.//0.00037:198:31//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P14737
- 35 F-NT2RM4001594//IMMEDIATE-EARLY PROTEIN IE180.//1.9e-05:147:34//PSEUDORABIES VIRUS (STRAIN KAPLAN) (PRV),//P33479
  - F-NT2RM4001597//THIOL:DISULFIDE INTERCHANGE PROTEIN TLPA (CYTOCHROME C BIOGENESIS PRO-TEIN TLPA).//5.7e-06:122:29//BRADYRHIZOBIUM JAPONICUM.//P43221
- F-NT2RM4001605//NUCLEAR PORE COMPLEX PROTEIN NUP155 (NUCLEOPORIN NUP155) (155 KD NU-40 CLEOPORIN) (P140).//1.7e-128:249:96//RATTUS NORVEGICUS (RAT).//P37199
- F-NT2RM4001611//SIS2 PROTEIN (HALOTOLERANCE PROTEIN HAL3),//1.5e-35:128:47//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST),//P36024
  - F-NT2RM4001629//MAGUK P55 SUBFAMILY MEMBER 3 (MPP3 PROTEIN) (DISCS, LARGE HOMOLOG 3),// 5.8e-42:254:37//HOMO SAPIENS (HUMAN).//Q13368
- 45 F-NT2RM4001650//HOMEOBOX PROTEIN HOX-A4 (CHOX-1.4).//0.62:19:57//GALLUS GALLUS (CHICKEN).//
  - F-NT2RM4001662//PROTEIN KINASE C, ALPHA TYPE (EC 2.7.1.-) (PKC-ALPHA).//0.29:90:32//HOMO SAPI-ENS (HUMAN),//P17252
  - F-NT2RM4001666//HYPOTHETICAL 48.6 KD PROTEIN IN ALPA-GABP INTERGENIC REGION. //1.1e-31:137: 44//ESCHERICHIA COLL//P37339
  - F-NT2RM4001682//PROBABLE 60S RIBOSOMAL PROTEIN L22.//0.98:55:29//CAENORHABDITIS ELEGANS.// P52819
    - F-NT2RM4001710//HYPOTHETICAL PROTEIN KIAA0039 (FRAGMENT), //0.56:113:28//HOMO SAPIENS (HU-MAN).//Q15054
- F-NT2RM4001714//SEPTIN 2 HOMOLOG (FRAGMENT).//1.4e-108:255:77//HOMO SAPIENS (HUMAN).// Q14141
  - F-NT2RM4001715//HYPOTHETICAL PROTEIN C19G10.16 IN CHROMOSOME I (FRAGMENT).//2.1e-36:148: 38//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10342

- F-NT2RM4001731//HYPOTHETICAL 54.9 KD PROTEIN C02F5.7 IN CHROMOSOME III.//1.1e-05:90:33// CAENORHABDITIS ELEGANS //P34284
- F-NT2RM4001741//TALIN.//1.1e-106:208:99//MUS MUSCULUS (MOUSE).//P26039
- F-NT2RM4001746//EBNA-1 NUCLEAR PROTEIN.//1.6e-09:155:38//EPSTEIN-BARR VIRUS (STRAIN B95-8)
- F-NT2RM4001754//COLLAGEN ALPHA 5(IV) CHAIN PRECURSOR.//0.93:158:33//HOMO SAPIENS (HUMAN).// P29400
  - F-NT2RM4001758//PUTATIVE SERINE/THREONINE-PROTEIN KINASE P78 (EC 2.7.1.-).//5.1e-113:277:79//
    HOMO SAPIENS (HUMAN).//P27448
- 10 F-NT2RM4001776/MYOSIN I ALPHA (MMI-ALPHA),//2 2e-73:262:54/MUS MUSCULUS (MOUSE),//P46735 F-NT2RM4001783//ZINC FINGER PROTEIN HRX (ALL-1) (FRAGMENT),//5.3e-26:169:39/MUS MUSCULUS (MOUSE),//P55200
  - F-NT2RM4001810//MALE SPECIFIC SPERM PROTEIN MST84DB.//2.3e-05:68:42//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643
- 15 F-NT2RM4001813//RHODOCETIN ALPHA SUBUNIT.//2.3e-05:115:34//AGKISTRODON RHODOSTOMA (MA-LAYAN PIT VIPER) (CALLOSELASMA RHODOSTOMA).//P81397
  - F-NT2RM4001819//CELL SURFACE GLYCOPROTEIN EMR1 PRECURSOR (EMR1 HORMONE RECEPTOR) (CELL SURFACE GLYCOPROTEIN F4/80).//1.7e-06:159:25//MUS MUSCULUS (MOUSE).//Q61549
  - F-NT2RM4001823//ZINC FINGER PROTEIN ZIC1 (ZINC FINGER PROTEIN OF THE CEREBELLUM 1).//2.6e-18:114:40//MUS MUSCULUS (MOUSE).//P46684
    - F-NT2RM4001828//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2).//4.0e-81:253:59//HOMO SA-PIENS (HUMAN).//P51523
    - F-NT2RM4001836//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT).//0.21:176:30//NEPHILA CLA-VIPES (ORB SPIDER).//P46804
- 25 F-NT2RM4001841//PROLINE-RICH PEPTIDE P-B.//0.046:27:40//HOMO SAPIENS (HUMAN).//P02814 F-NT2RM4001842//HYPOTHETICAL 7.0 KD PROTEIN B03B8.1 IN CHROMOSOME III.//0.98:35:42//
  - F-NT2RM4001842/HYPOTHETICAL 7.0 KD PROTEIN B03B8.1 IN CHROMOSOME III.//0.98:35:42// CAENORHABDITIS ELEGANS.//Q11104
  - F-NT2RM4001856/HYPOTHETICAL 75.2 KD PROTEIN IN ACS1-GCV3 INTERGENIC REGION.//2.3e-37:242: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39722
- 30 F-NT2RM4001858/T-BOX PROTEIN VEGT (T-BOX PROTEIN BRAT) (T-BOX PROTEIN ANTIPODEAN).//1.8e-23:78:64//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P87377
  - F-NT2RM4001865//NEURONAL CALCIUM SENSOR 2 (NCS-2).//0.012:83:28//CAENORHABDITIS ELEGANS.// P36609
  - F-NT2RM4001876//HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECUR-SOR.//3.8e-10:242:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47179
    - F-NT2RM4001880/EC PROTEIN HOMOLOG./I0.22:59:32//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//
      P93746
      F-NT2RM4001905//60S RIBOSOMAL PROTEIN L40 (CEP52).//0.57:20:60//HOMO SAPIENS (HUMAN), RAT-
    - TUS NORVEGICUS (RAT), AND GALLUS GALLUS (CHICKEN).//P14793
- 40 F-NT2RM4001922

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- F-NT2RM4001930//PUTATIVE GLUCOSYLTRANSFERASE C08B11.8 (EC 2.4.1.-).//5.5e-45:167:53// CAENORHABDITIS ELEGANS.//Q09226
- F-NT2RM4001938//RTOA PROTEIN (RATIO-A).//0.0036:120:32//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P54681
- 45 F-NT2RM4001940/IROQUOIS-CLASS HOMEODOMAIN PROTEIN IRX-1 (FRAGMENT)//0.32:31:48//HOMO SAPIENS (HUMAN).//P78415
  - F-NT2RM4001953//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//2.2e-43:56:85//HOMO SAPIENS (HUMAN).// P39192
  - F-NT2RM4001965//IG ALPHA-1 CHAIN C REGION.//0.56:73:34//GORILLA GORILLA GORILLA (LOWLAND GORILLA).//P20758
  - F-NT2RM4001969//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONES CP3, CP4 AND CP5) [CON-TAINS: BASIC PEPTIDE IB-6: PEPTIDE P-HI //0.0016:140:27/HOMO SAPIENS (HUMAN) //P04280
    - F-NT2RM4001979//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2).//3.9e-21:103:51//HOMO SA-PIENS (HUMAN).//P51523
- 55 F-NT2RM4001984//HYPOTHETICAL PROTEIN LAMBDA-SP5.//0.0034;50:40//MUS MUSCULUS (MOUSE).// P15974
  - F-NT2RM4001987//IRREGULAR CHIASM C-ROUGHEST PROTEIN PRECURSOR (IRREC PROTEIN).//6.9e-17:115:31//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q08180

- F-NT2RM4002013//HYPOTHETICAL 54.5 KD TRP-ASP REPEATS CONTAINING PROTEIN ZC302.2 IN CHRO-MOSOME V //0.0062:117:28//CAENORHARDITIS ELEGANS //023256
- F-NT2RM4002018//SPORE COAT PROTEIN SP96.//4.3e-06:203:28//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P14328
- F-NT2RM4002034/RHO-GAP HEMATOPOIETIC PROTEIN C1 (P115) (KIAA0131).//0.78:132:25//HOMO SAPIENS (HUMAN).//P98171
  - F-NTZRM4002044/VITELLOGENIN I PRECURSOR (MINOR VITELLOGENIN) [CONTAINS: LIPOVITELLIN I (LVI); PHOSVITIN (PV); LIPOVITELIN II (LVI); YGP42]//10.082/21/24//GALLUS GALLUS (CHICKEN)//P87498 F-NTZRM4002054//DUPLICATE PROCYCLIN/10/007944-55///RYPANOSOMA BRUCEI BRUCEI.//P14047
- f-NT2RM4002056//PUTATIVE Z PROTEIN./IO.82:39:30//OVIS ARIES (SHEEP).//P08105 F-NT2RM4002062//ASPARTYL-TRNA SYNTHETASE (EC 6.1.1.12) (ASPARTATE.-TRNA LIGASE) (ASPRS).// 7.0e-37:80:52//THERMUS AQUATICUS (SUBSE THERMOPHILUS).//P36419
  - F-NT2RM4002063//SARCOSINE OXIDASE (EC 1.5.3.1)//2.2e-25:216:31//BACILLUS SP. (STRAIN NS-129).// P23342
- 15 F-NT2RM4002066//HYPOTHETICAL PROTEIN KIAA0192 (FRAGMENT),//1.1e-94:260:71//HOMO SAPIENS (HUMAN),//Q93074
  - F-NT2RM4002067//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.5e-15:51:70//HOMO SAPIENS (HUMAN).//
- F-NT2RM4002073//ELASTIN PRECURSOR (TROPOELASTIN).//4.9e-05:88:36//HOMO SAPIENS (HUMAN).//
  - F-NT2RM4002075//RING CANAL PROTEIN (KELCH PROTEIN).//7.2e-43:220:41//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
    - F-NT2RM4002093/POLYPYRIMIDINE TRACT-BINDING PROTEIN (PTB) (HETEROGENEOUS NUCLEAR RI-BONUCLEOPROTEIN I) (HNRNP I) (57 KD RNA-BINDING PROTEIN PPTB-1).//1.8e-93.255:72//HOMO SAPI-ENS (HUMAN)./P26599
    - F-NT2RM4002109/IKINESIN-LIKE PROTEIN KIF4.//3.7e-101:260:78//MUS MUSCULUS (MOUSE).//P33174 F-NT2RM4002128//HYPOTHETICAL PROTEIN IN CYCB 3'REGION PRECURSOR (ORF2) (FRAGMENT).// 0 91:49:32/PARACOCCUS DENITRIFICANS //P29969
  - F-NT2RM4002140//GROUCHO PROTEIN (ENHANCER OF SPLIT M9/10).//0.36:104:22//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P16371
- F-NT2RM4002145//SLIT PROTEIN PRECURSOR.//8.6e-13:127:33//DROSOPHILA MELANOGASTER (FRUIT FLY) //P24014
  - F-NT2RM4002146//MAGO NASHI PROTEIN.//7.9e-69:143:91//DROSOPHILA MELANOGASTER (FRUIT FLY).//
- 35 F-NT2RM4002161//DUAL SPECIFICITY PROTEIN PHOSPHATASE (EC 3.1.3.48) (EC 3.1.3.16) // 0.0062:99:26// CHLAMYDOMONAS EUGAMETOS.//Q39491
  - F-NT2RM4002174//MRP PROTEIN.//4.5e-50:183:55//ESCHERICHIA COLL//P21590

- F-NT2RM4002189//MUCIN 2 PRECURSOR (INTESTINAL MUCIN 2).//2.6e-14:233:29//HOMO SAPIENS (HU-MAN).//Q02817
- 40 F-NT2RM4002194/TRANSMEMBRANE PROTEIN SEX PRECURSOR.//0.92:108:28//HOMO SAPIENS (HU-MAN).//P51805
  - F-NT2RM4002205//ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR (MEF-G).//5.8e-39:122:72// RATTUS NORVEGICUS (RAT).//Q07803
- F-NT2RM4002213//HYPOTHETICAL 88.4 KD PROTEIN B0464.7 IN CHROMOSOME III.//9.9e-27:110:43//
  45 CAENORHABDITIS ELEGANS.//Q03565
  - F-NT2RM4002226//GTPASE ACTIVATING PROTEIN ROTUND.//1.3e-21:147:41//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P40809
    - F-NT2RM4002251//PROTEIN EF-7 (FRAGMENT).//0.00082:45:42//MUS MUSCULUS (MOUSE).//P97805
- F-NT2RM4002256//COLD-REGULATED PROTEIN 1 (FRAGMENT)///0.00015:114:42//HORDEUM VULGARE (BARLEY)//P23251
  F-NT2RM4002256//CUTICLE COLLAGEN 2//0.00013:142:33//CAENORHABDITIS ELEGANS//P17656
- F-NTZRM-002278/INTPOTHETICAL 22.2 KD PROTEIN IN NSR1-TIF4631 INTERGENIC REGION.//1.0-40:52// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53288 E-NTZPM-002278/INTPOTHETICAL 22.2 KD PROTEIN IN NSR1-TIF4631 INTERGENIC REGION.//1.0-40:52// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53288
- 55 F-NT2RM4002287//GAR2 PROTEIN://0.00055:225:23//SCHIZOSACCHAROMYCES POMBE (FISSION YFAST) //P41891
  - F-NT2RM4002294//HYPOTHETICAL PROTEIN KIAA0281 (HA6725).//1.1e-60:152:75//HOMO SAPIENS (HU-MAN).//Q92556

- F-NT2RM4002301//GENERAL STRESS PROTEIN CTC (FRAGMENT).//0.56:43:39//BACILLUS CALDOLYTI-CUS //P42832
- F-NT2RM4002323//NONHISTONE CHROMOSOMAL PROTEIN HMG-17.//0.0080;73:35//BOS TAURUS (BO-VINE).//P02313
- F-NT2RM4002339//METALLOTHIONEIN 10-III (MT-10-III).//0.67:34:38//MYTILUS EDULIS (BLUE MUSSEL).//
  - F-NT2RM4002344//METALLOTHIONEIN-I (MT-I).//0.84:41:31//MUS MUSCULUS (MOUSE).//P02802
  - F-NT2RM4002373//GLUTENIN, HIGH MOLECULAR WEIGHT SUBUNIT DY10 PRECURSOR.//0.0019:190:28//TRITICUM AESTIVUM (WHEAT).//P10387
- 10 F-NT2RM4002374//5E5 ANTIGEN.//0.0059:170:32//RATTUS NORVEGICUS (RAT).//Q63003
  - F-NT2RM4002383//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.13:17:88//HOMO SAPIENS (HUMAN),//
  - F-NT2RM4002390
  - F-NT2RM4002398//HNRNP ARGININE N-METHYLTRANSFERASE (EC 2.1.1.-) (ODP1 PROTEIN).//0.034:110: 27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38074
- 15 27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),/IP38074 F-NT2RM4002409//ACETYL-COENZYME A SYNTHETASE (EC 6.2.1.1) (ACETATE--COA LIGASE) (ACYL-AC-TIVATING ENZYME)./I4 (be-20.179:31/METHANOTHRIX SOEHNGENII./IP27095
  - F-NT2RM4002438//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.7e-15:41:95//HOMO SAPIENS (HUMAN).//
- F-NT2RM4002446//CRYPTDIN-RELATED PROTEIN 4C-1 PRECURSOR (CRS4C).//0.0058:24:50//MUS MUS-CULUS (MOUSE).//P17534
  - F-NT2RM4002452//METALLOTHIONEIN 10-II (MT-10-II).//0.83:48:37//MYTILUS EDULIS (BLUE MUSSEL).//
- F-NT2RM4002457//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//4.9e-07:52:63//HOMO SAPIENS (HUMAN).//
- 25 P39192 F-NT2RM4002460//C-HORDEIN (CLONE PC-919) (FRAGMENT).//0.92:43:30//HORDEUM VULGARE (BAR-
  - LEY).//P17992
    F-NT2RM4002479//RNA HELICASE-LIKE PROTEIN DB10.//1.7e-28:200:41//NICOTIANA SYLVESTRIS (WOOD
- TOBACCO)./IP46942

  F-NT2RM4002482/IH/POTHETICAL 65.9 KD PROTEIN YPR065W.//8.8e-26:123:49//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST)./IO12514 F-NT2RW4002493/IJ.ARVAL CUTICLE PROTEIN | PRECURSOR //IO.17:
  - 126:27//DROSOPHILA MIRANDA (FRUIT FLY).//P91627 F-NTZRM4002499//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/9.4e-34:92:80//HOMO SAPIENS (HUMAN).// P30194
- 35 F-NT2RM4002504//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//3.4e-19:55:83//HOMO SAPIENS (HUMAN).//
  - F-NT2RM4002527//WD-40 REPEAT PROTEIN MSI2.//3.0e-07:193:27//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//O22468
  - F-NT2RM4002532//AEROLYSIN REGULATORY PROTEIN.//0.97:19:47//AEROMONAS SOBRIA.//P09165
- 40 F-NT2RM4002534/MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L32 PRECURSOR (YML32).//0.76:86:22// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P25348
  - F-NT2RM4002558/ILONG-CHAIN FATTY ACID TRANSPORT PROTEIN (FATP) J/4.2e-55:204:50//MUS MUSCULUS (MOUSE).//Q60714
- F-NT2RM4002565//CHYMOTRYPSIN/ELASTASE ISOINHIBITORS 2 TO 5.//1.0:16:62//ASCARIS SUUM (PIG ROUNDWORM) (ASCARIS LUMBRICOIDES).//P07852
- F-NT2RM4002567//HYPOTHETICAL 74.0 KD PROTEIN IN CAJ1-HOM3 INTERGENIC REGION.//2.7e-10:184: 29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40032
  - F-NT2RM4002571//POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (EC 2.4.1.41) (PROTEIN-UDP ACETYLGALACTOSAMINYLTRANSFERASE) (UDP-GALNAC:POLYPEPTIDE, N- ACETYLGALACTOS-
- 40 AMINYLTRANSFERASE) (GALNAC-T1).//2.4e-25:124:47//HOMO SAPIENS (HUMAN).//Q10472 F-NT2RM4002593//HYPOTHETICAL 9.1 KD PROTEIN IN TETB-EXOA INTERGENIC REGION.//0.95:36:38//BA-
  - CILLUS SUBTILIS.//P37509
    F-NT2RM4002594//MSP1 PROTEIN HOMOLOG.//9.0e-68:227:60//CAENORHABDITIS ELEGANS.//P54815
- F-NT2RM4002623//ASPARTYL-TRNA SYNTHETASE (EC 6.1.1.12) (ASPARTATE--TRNA LIGASE) (ASPRS).//
  3.3e-54-243-47/JSYNECHOCYSTIS SP. (STRAIN PCC 6803) //P73851
  - F-NT2RP1000018//SUPPRESSOR PROTEIN SRP40.//0.0023:131:25//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583
    - F-NT2RP1000035//RING CANAL PROTEIN (KELCH PROTEIN).//1.0e-06:63:34//DROSOPHILA MELA-

#### NOGASTER (FRUIT FLY).//Q04652

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F-NT2RP1000040//LETHAL NEUROTOXIN TX1.//0.69:21:47//PHONEUTRIA NIGRIVENTER (BRAZILIAN ARMED SPIDER).//P17727

F-NT2RP1000063//HYPOTHETICAL 25.1 KD PROTEIN IN SMC3-MRPL8 INTERGENIC REGION.//3.8e-14:130: 30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40359

F-NT2RP1000086//HYPOTHETICAL 9.4 KD PROTEIN IN RNPA-THDF INTERGENIC REGION.//0.16:44:40//ES-CHERICHIA COLL//P22847

F-NT2RP1000101//45.8 KD PROTEIN IN SHM1-MRPL37 INTERGENIC REGION.//1.9e-06:74:32//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P38344

10 F-NT2RP1000111/I/COP1 REGULATORY PROTEIN (FUSCA PROTEIN FUS1).//2.7e-19:135:36//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P43254

F-NT2RP1000112//DUAL SPECIFICITY PROTEIN KINASE TTK (EC 2.7.1.-) (PYT).//1.2e-39:91:62//HOMO SA-PIENS (HUMAN).//P33981

F-NT2RP1000124//ATP-DEPENDENT PROTEASE LA 2 (EC 3.4.21.53).//0.074:131:24//MYXOCOCCUS XAN-THUS //P36774

F-NT2RP1000130//HEPATOMA-DERIVED GROWTH FACTOR (HDGF).//1.5e-49:186:56//MUS MUSCULUS (MOUSE).//P51859

F-NT2RP1000163//METALLOTHIONEIN (MT).//0.98:41:34//PLEURONECTES PLATESSA (PLAICE).//P07216 F-NT2RP1000170//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-

MENT)//0.85:64:35/H-OMO SAPIENS (HUMAN).//P10162 F-NT2RP100174///MMEDIATE-EARLY PROTEIN IE180.//0.00056:89:37//PSEUDORABIES VIRUS (STRAIN KAPLAN) (PRV).//P33479

F-NT2RP1000191//NIFU PROTEIN.//0.53:78:35//FRANKIA ALNI.//P46045

F-NT2RP1000202//ANKYRIN, BRAIN VARIANT 2 (ANKYRIN B) (ANKYRIN, NONERYTHROID) (FRAGMENT).//
9.1e-21:148:39//HOMO SAPIENS (HUMAN).//Q01485

F-NT2RP1000243//HYPOTHETICAL PROTEIN MJ1136.//1.4e-37:219:36//METHANOCOCCUS JANNASCHII.// Q58536

F-NT2RP1000259//HYPOTHETICAL PROTEIN TP0318.//0.18:25:44//TREPONEMA PALLIDUM.//083338 F-NT2RP1000272//SPLICING FACTOR, ARGININE/SERINE-RICH 3 (PRE-MRNA SPLICING FACTOR SRP20)

 (X16 PROTEIN)/II. 6e-18:133:36/HOMO SAPIENS (HUMAN), AND MUS MUSCULUS (MOUSE).//P23152 F-NT2RP1000324

F-NT2RP1000326//HYPOTHETICAL 29.8 KD PROTEIN ZC97.1 IN CHROMOSOME III.//1.0e-23:129:36// CAENORHABDITIS ELEGANS.//P34599

F-NT2RP1000333//ANTI-SILENCING PROTEIN 1.//2.5e-45:147:57//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32447

F-NT2RP1000348//REDUCED VIABILITY UPON STARVATION PROTEIN 161.//4.8e-14:119:34//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P25343

F-NT2RP1000357//TRYPOMASTIGOTE DECAY-ACCELERATING FACTOR (T-DAF) (FRAGMENT).//1.0:43:32//TRYPANOSOMA CRUZI.//Q26327

40 F-NT2RP1000358//HYPOTHETICAL 84.4 KD PROTEIN IN RPC2/RET1 3'REGION.//7.9e-28:244:35//SACCHA-ROMYCES CEREVISIAE (BAKER'S YEAST).//P39744

F-NT2RP1000363//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//2.2e-07:178:30// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437

F-NT2RP1000376//ANKYRIN R (ANKYRINS 2.1 AND 2.2) (ERYTHROCYTE ANKYRIN).//1.5e-20:254:31//HOMO SAPIENS (HUMAN).//P16157

F-NT2RP1000409//CYTOCHROME C3 (CYTOCHROME C7) (C551.5),//1.0:34:26//DESULFUROMONAS ACE-TOXIDANS (CHLOROPSEUDOMONAS ETHYLICA),//P00137

F-NT2RP1000413//MEMBRANE-ASSOCIATED PROTEIN HEM-2 (NAP1 PROTEIN).//3.7e-131:230:97//RAT-TUS NORVEGICUS (RAT).//P55161

F-NT2RP1000416//SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.83:54:40//DROSOPHILA SIMULANS (FRUIT FLY).//P13729

F-NT2RP1000418//HYPOTHETICAL 9.9 KD PROTEIN IN GCVT-SPOIIIAA INTERGENIC REGION.//0.24:91:35// BACILLUS SUBTILIS.//P49779

F-NT2RP1000439//HYPOTHETICAL 100.5 KD PROTEIN C1B9.04 IN CHROMOSOME I.//0.13:172:22// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10429

F-NT2RP1000443//QUINONE OXIDOREDUCTASE (EC 1.6.5.5) (NADPH:QUINONE REDUCTASE) (ZETA-CRYSTALLIN).//1.9e-08:167:24//HOMO SAPIENS (HUMAN).//Q08257

F-NT2RP1000460//NUCLEAR MOVEMENT PROTEIN NUDC.//1.0e-18:149:34//EMERICELLA NIDULANS (AS-

PERGILLUS NIDULANS) //P17624

US AETHIOPS (GREEN MONKEY) (GRIVET).//P33194

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- F-NT2RP1000470//PUTATIVE ATP-DEPENDENT RNA HELICASE T26G10.1 IN CHROMOSOME III.//1.3e-43: 180:47//CAENORHABDITIS ELEGANS.//P34580
- F-NT2RP1000478//TUBULIN BETA-6 CHAIN (CLASS-VI).//1.5e-45:85:63//GALLUS GALLUS (CHICKEN).// P09207
- F-NT2RP1000481//HYPOTHETICAL 5.8 KD PROTEIN IN PUHA 5'REGION (ORF55).//0.083:21:47//RHODO-BACTER CAPSULATUS (RHODOPSEUDOMONAS CAPSULATA).//P26159
- F-NT2RP1000493//POSSIBLE DNA-REPAIR PROTEIN XP-E (POSSIBLE XERODERMA PIGMENTOSUM GROUP E PROTEIN) (UV-DAMAGED DNA-BINDING PROTEIN) (UV-DDB).//6.6e-11:139:31//CERCOPITHEC-
- F-NT2RP1000513//60S RIBOSOMAL PROTEIN L22.//0.017:92:30//DROSOPHILA MELANOGASTER (FRUIT FLY).//P50887
- F-NÍZRP1000522//UBIQUITIN CARBOXYL-TERMINAL HYDROLASE DUB-1 (EC 3.1.2.15) (UBIQUITIN THI-OLESTERASE DUB-1) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE DUB-1) (DEUBIQUITINATING EN-ZYME 11/0.0058:883/MUS MUSCULUS (MOUSE)//Q61088
- F-NT2RP1000547//COP-COATED VESICLE MEMBRANE PROTEIN P24 PRECURSOR (FRAGMENT).//1.2e-09:69:36//CRICETULUS GRISEUS (CHINESE HAMSTER).//P49020
  - F-NT2RP1000574//HOMEOBOX PROTEIN MEIS2 (MEIS1-RELATED PROTEIN 1).//6.0e-39:141:65//MUS MUS-CULUS (MOUSE).//P97367
- 20 F-NT2RP1000577//PUTATIVE ATP-DEPENDENT RNA HELICASE YDL031W.//0.00016:48:45//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//Q12389
- F-NT2RP1000581//VON WILLEBRAND FACTOR PRECURSOR.//0.00017:61:50//HOMO SAPIENS (HUMAN).//
- F-NT2RP1000609//LINOLEOYL-COA DESATURASE (EC 1.14.99.25) (DELTA(6)-DESATURASE).//4.4e-07:128: 31//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//Q08871
  - F-NT2RP1000629//CLATHRIN COAT ASSEMBLY PROTEIN AP47 (CLATHRIN COAT ASSOCIATED PROTEIN AP47) (GOLGI ADAPTOR AP-1 47 KD PROTEIN) (HA1 47 KD SUBUNIT) (CLATHRIN ASSEMBLY PROTEIN ASSEMBLY PROTEIN COMPLEX 1 MEDIUM CHAIN),//4 2e-70-167-86/MUS MUSCULUS (MOUSE),///735585 F-NT2RP1000630/HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DALS INTERGENIC REGION PRECURSOR.//
- 30 0.0011:238:21//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47179 F-NT2RP1000677//COLLAGEN ALPHA 1(XVI) CHAIN PRECURSOR.//0.99:71:33//HOMO SAPIENS (HUMAN).// O07/092
  - F-NT2RP1000688//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.0024:19:94//HOMO SAPIENS (HUMAN).//
- 35 F-NT2RP1000695//HYPOTHETICAL 83.8 KD PROTEIN C27F2.7 IN CHROMOSOME III.//2.2e-30:185:37// CAENORHABDITIS ELEGANS.//Q18262
  - F-NT2RP1000701//PHOSPHOLIPASE A-2-ACTIVATING PROTEIN (PLAP).//3.2e-65:128:93//RATTUS NOR-VEGICUS (RAT) //P54319
- F-NT2RP1000721//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII-135) (TAFII-130) (TAFII-130) //2 3e-06:139:34//HOMO SAPIENS (HUMAN) //000268
  - F-NT2RP1000730//MYOSIN LIGHT CHAIN 1, SLOW-TWITCH MUSCLE B/VENTRICULAR ISOFORM (FRAG-MENT).//0.89:40:40//MUS MUSCULUS (MOUSE).//P09542
    - F-NT2RP1000733//METALLOTHIONEIN-LIKE PROTEIN CRS5.//0.024:24:45//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P41902
- 45 F-NT2RP1000738//SALIVARY ACIDIC PROLINE-RICH PHOSPHOPROTEIN 1/2 PRECURSOR (PRP-1 / PRP-3) (PRP-2 / PRP-4) (PIF-5 ) (PROTEIN A / PROTEIN C) [CONTAINS: PEPTIDE P-C].//0.040:82:36//HOMO SAPIENS (HUMAN)./P02810
  - F-NT2RP1000746//HYPOTHETICAL 27.1 KD PROTEIN UFD4-CAP1 INTERGENIC REGION.//2.0e-30:170:37// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P33201
  - F-NT2RP1000767//PSEUDOMONAPEPSIN PRECURSOR (EC 3.4.23.37) (PEPSTATIN-INSENSITIVE CAR-BOXYL PROTEINASE)//0.99:75:34//PSEUDOMONAS SP. (STRAIN 101).//P42790
    - F-NTZRP1000782/CELL SURFACE GLYCOPROTEIN A15 (T-CELL ACUTE LYMPHOBLASTIC LEUKEMIA AS-SOLIATED ANTIGEN 1) (TALLA-1) (MEMBRANE COMPONENT, X CHROMOSOME, SURFACE MARKER 1).// 23-e23:159:35/HONO SAPIENS (HUMAN).//P41732
- 55 F-NT2RP1000796//CORNIFIN (SMALL PROLINE-RICH PROTEIN I) (SPR-I) (SMALL PROLINE-RICH SQUA-MOUS CELL MARKER) (SPRP).//0.00018:79:32//SUS SCROFA (PIG).//P35323
  - F-NT2RP1000825//GTPASE-ACTIVATING PROTEIN RHOGAP (RHO-RELATED SMALL GTPASE PROTEIN ACTIVATOR) (CDC42 GTPASE-ACTIVATING PROTEIN) (P50-RHOGAP).//3.1e-37:89:64//HOMO SAPIENS (HU-

MAN).//Q07960

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- F-NT2RP1000833//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.32:29:48//HOMO SAPIENS (HLIMAN) //P22531
- F-NT2RP1000834//2-ARYLPROPIONYL-COA EPIMERASE (EC 5.-.-).//6.4e-67:202:68//RATTUS NORVEGI-CUS (RAT) //P70473
  - F-NT2RP1000836//HYPOTHETICAL 7.3 KD PROTEIN IN 100 KD PROTEIN REGION.//1.0:35:54//HUMAN AD-ENOVIRUS TYPE 41.//P23691
- F-NT2RP1000846//SMALL PROLINE-RICH PROTEIN 2-1.//0.013:35:48//HOMO SAPIENS (HUMAN).//P35326 F-NT2RP1000851//PERIOD CLOCK PROTEIN (FRAGMENT).//0.082:28:57//DROSOPHILA SALTANS (FRUIT
- F-NT2RP1000851//PERIOD CLOCK PROTEIN (FRAGMENT).//0.082:28:57//DROSOPHILA SALTANS (FRUI FLY).//Q04536
  - F-NT2RP1000856//PLATELET-ENDOTHELIAL TETRASPAN ANTIGEN 3 (PETA-3) (GP27) (MEMBRANE GLYC-OPROTEIN SFA-1) (CD151 ANTIGEN).//2.5e-26:190:30//MUS MUSCULUS (MOUSE).//035566
- F-NT2RP1000860//POTENTIAL TRANSCRIPTIONAL ADAPTOR.//0.13:86:36//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//Q02336
- F-NT2RP1000902//HYPOTHETICAL 127.4 KD PROTEIN F07F6.4 IN CHROMOSOME III.//7.6e-11:200:35// CAENORHABDITIS ELEGANS.//Q09531
  - F-NTZRP1000315/HYPOTHETICAL GTP-BINDING PROTEIN IN PMI40-PAC2 INTERGENIC REGION.//1.4e-06-88:35/ISACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40010 F-NTZRP1000316//SUPPRESSOR PROTEIN SRP40//0.40:00:35/ISACCHAROMYCES CEREVISIAE (BAK-
  - ER'S YEAST).//P32583
  - F-NT2RP1000943//MUCIN 2 PRECURSOR (INTESTINAL MUCIN 2).//0.099:75:34//HOMO SAPIENS (HU-MAN).//Q02817
    - F-NT2RP1000944//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//7.6e-06:65:41//ORGYIA PSEUDOTSUGATA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//O10341
- 22 F-NT2RP1000947/JUBIGUITIN-CONJUGATING ENZYME E2-17 KD 2 (EC 6.3.2.19) (JUBIGUITIN- PROTEIN LIGASE) (JUBIGUITIN CARRIER PROTEIN) (E2(17)MB 2)/3.66-12-2777/H/OMO SAPIENS (HUMAN), MUS MUSCULUS (MOUSE), RATTUS NORVEGICUS (RAT), AND XENOPUS LAEVIS (AFRICAN CLAWED FROG).// PR1889
  - F-NT2RP1000954//RING CANAL PROTEIN (KELCH PROTEIN).//2.8e-15:169:28//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//004652
- F-NT2RP1000958//HYPOTHETICAL GTP-BINDING PROTEIN IN PMI40-PAC2 INTERGENIC REGION.//4.2e-16:162:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40010
  - F-NT2RP1000959//CORNIFIN A (SMALL PROLINE-RICH PROTEIN IA) (SPR-IA) (SPRK).//0.0031:34:44//HOMO SAPIENS (HUMAN).//P35321
- 55 F-NT2RP1000986/INUCLEOLIN (PROTEIN C23)//1.5e-52:110:95//HOMO SAPIENS (HUMAN)/P19338 F-NT2RP1000986/INUGHT-HARVESTING PROTEIN B-1015, LPHA CHAIN PRECURSOR (ANTENNA PIG-MENT PROTEIN, ALPHA CHAIN)//0.87:37:45//RHODOPSEUDOMONAS VIRIDIS://P04123 F-NT2RP1000988
  - F-NT2RP1001011//PROTEIN P19.//0.96:30:50//BACTERIOPHAGE PRD1.//P17638
- 40 F-NT2RP1001013//DNA-BINDING PROTEIN 65 (PROTEIN GP65).//1.0:20:45//BACTERIOPHAGE T4.//P16012 F-NT2RP1001014
  - F-NT2RP1001033//TUBULIN GAMMA CHAIN.//2.5e-16:112:42//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P25295
- F-NT2RP1001073//HYPOTHETICAL 10.4 KD PROTEIN IN FTR1-SPT15 INTERGENIC REGION.//7.6e-16:82: 51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40089
  - F-NT2RP1001079//SARCOSINE OXIDASE (EC 1.5.3.1).//4.8e-15:95:40//ARTHROBACTER SP. (STRAIN TE1826).//P40873
    - F-NT2RP1001080//PROBABLE ATP-DEPENDENT RNA HELICASE DBP9.//2.4e-29:126:46//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//Q06218
- 50 F-NT2RP1001113//SMALL PROLINE-RICH PROTEIN 2-1/10, 49:38:39//HOMO SAPIENS (HUMAN).//P35326 F-NT2RP1001173//RHOMBOTIN-1 (CYSTEINE RICH PROTEIN TTG-1) (T-CELL TRANSLOCATION PROTEIN 11/LM-ONLY PROTEIN 11/L0)95:54:37//HOMO SAPIENS (HUMAN)//P25800
  - F-NT2RP1001177//HISTONE MACRO-H2A.1.//1.6e-29:85:76//RATTUS NORVEGICUS (RAT).//Q02874
- 55 F-NT2RP1001199//NEUROTOXIN I.//1.0:23:47//CENTRUROIDES SCULPTURATUS (BARK SCORPION).// P01491
  - F-NT2RP1001247//TRANSFORMING GROWTH FACTOR BETA 4 PRECURSOR (TGF-BETA 4) (ENDOMETRI-AL BLEEDING-ASSOCIATED FACTOR).//3.3e-08:28:89//HOMO SAPIENS (HUMAN).//000292

- F-NT2RP1001248/tAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN).//0.33:49:28//HUMAN IMMU-NODEFICIENCY VIRUS TYPE 1 (NDK ISOLATE) (HIV-1).//P18804
  F-NT2RP1001253//GLUCOSAMINE-6-PHOSPHATE ISOMERASE (EC 5.3.1.10) (GLUCOSAMINE-6-PHOS-
- PHATE DEAMINASE) (GNPDA) (OSCILLIN) (KIAAO080) //3.8e-46.115.81/IHOMO SAPIENS (HUMAN) //F46926 F-NT2RP1001288/GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PRO-TEIN) (35 KD LECTIN) (CARBOHYDRATE BINDING PROTEIN 35) (CBP 35) (LAMININ-BINDING PROTEIN) (LECTIN L-29) (L-34 GALACTOSIDE-BINDING LECTIN) //IO.16.48:37/MUS MUSCULUS (MOUSE) //IP16110 F-NT2RP1001294/MICROTUBLIC-ASSOCIATED PROTEIN YTM1 //6.1e-05-92:34//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST) //O12024
- F-NT2RP1001302//MICROTUBULE-ASSOCIATED PROTEIN YTM1.//1.2e-05:92:33//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST).//Q12024
  - F-NTZRP1001310//PROBABLE E4 PROTEIN//0.99:109:26//HUMAN PAPILLOMAVIRUS TYPE 5.//P06924 F-NTZRP1001311//SODIUM/HYDROGEN EXCHANGER 5 (NA(+)H(+) EXCHANGER 5) (NHE-5) (FRAGMENT).//0.99:94:31//HOMO SAPIENS (HUMAN).//0.14940
- 15 F-NT2RP1001313//CYTOCHROME B5.//9.0e-13:92:38//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P40312
  - F-NT2RP1001361//NADH-UBIQUINONE OXIDOREDUCTASE SUBUNIT B14.5B (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-B14.5B) (CI-B14.5B).//1.2e-47:117:74//BOS TAURUS (BOVINE).//Q02827
  - F-NT2RP1001385//CELL DIVISION PROTEIN FTSN.//0.64:107:28//ESCHERICHIA COLI.//P29131
- F-NT2RP1001395//PROCOLLAGEN ALPHA 2(I) CHAIN PRECURSOR (FRAGMENTS).//0.25:35:45//GALLUS GALLUS (CHICKEN).//P02467
  - F-NT2RP1001410//PUTATIVE GTP-BINDING PROTEIN W08E3.3.//2.2e-41:129:67//CAENORHABDITIS ELE-GANS //P91917
- F-NT2RP1001424//UREASE ACCESSORY PROTEIN UREF (FRAGMENT).//0.87:24:45//ESCHERICHIA COLI.//
  25 003286
  - F-NT2RP1001432//CYSTEINE PROTEINASE INHIBITOR B (CYSTATIN B) (SCB).//1.0:35:42//HELIANTHUS ANNUUS (COMMON SUNFLOWER).//010993
    - F-NT2RP1001449//KERATIN, ULTRA HIGH-SULFUR MATRIX PROTEIN (UHS KERATIN).//0.053:37:37//OVIS ARIES (SHEEP) //P26372
- F-NTZRP1001457/IHYPOTHETICAL 57.0 KD TRP-ASP REPEATS CONTAINING PROTEIN IN CPR4-SSK22 INTERGENIC REGION //2.9e-16:159:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P25382 F-NTZRP1001466/IHYPOTHETICAL PROTEIN MJ0284 //5.3e-15:162:35//METHANOCOCCUS JANNASCHII.//
  - Q57732 F-NT2RP1001475//HYPOTHETICAL 195.1 KD PROTEIN IN DNA43-UBI1 INTERGENIC REGION.//0.69:119:27//
- 39 SACCHAROMYCES CEREVISIAE (BAKER'S YEAST), //Pa/du457 F-NT2RP1001482//PROTEASOME COMPONENT C9 (EC 3.4.99.46) (MACROPAIN SUBUNIT C9) (MULTICAT-ALYTIC ENDOPEPTIDASE COMPLEX SUBUNIT C9) //1 0:5832//HOMO SAPIENS (HUMAN) //P25789
  - F-NT2RP1001494//MALE STERILITY PROTEIN 2.//2.4e-12:84:42//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//Q08891
- 4º F-NT2RP1001543/MYO-INOSITOL-1-PHOSPHATE SYNTHASE (EC 5.5.1.4) (IPS).//6.3e-37:94:52//SPIRODE-LA POLYBRHIZA //P42803
  - F-NT2RP1001546//LEUKOCYTE SURFACE ANTIGEN CD53 (CELL SURFACE GLYCOPROTEIN CD53).//9.3e-11:98:29//HOMO SAPIENS (HUMAN).//P19397
- F-NT2RP1001569//SIGNAL RECOGNITION PARTICLE RECEPTOR BETA SUBUNIT (SR-BETA).//2.2e-64:159: 84//MUS MUSCULUS (MOUSE).//P47758
- F-NT2RP1001616//HYPOTHETICAL 13.5 KD PROTEIN C45G9.7 IN CHROMOSOME III.//9.2e-05:49:42//
  CAENORHABDITIS FLEGANS //O09606
  - F-NT2RP1001665//REGB PROTEIN.//0.99:29:37//PSEUDOMONAS AERUGINOSA.//Q03381

- F-NT2RP2000001//SMALL PROLINE-RICH PROTEIN 2-1 //10 64/36:41/IHOMO SAPIENS (HUMAN) //P35326 F-NT2RP20000006//DNAJ PROTEIN HOMOLOG 1 (HDJ-1) (HEAT SHOCK PROTEIN 40) (HSP40) //1.7e-19:74: 52/IHOMO SAPIENS (HUMAN) //P25685
  - F-NT2RP2000007//TROPOMYOSIN, FIBROBLAST AND EPITHELIAL MUSCLE-TYPE (TM36) (TME1) (TM1).// 0.93:126:23//HOMO SAPIENS (HUMAN).//P06468
- F-NT2RP2000008//ZINC FINGER PROTEIN 33A (ZINC FINGER PROTEIN KOX31) (KIAA0065) (HA0946)

  656 (FRAGMENT).//4 2e-35:156:54//HOMO SAPIENS (HUMAN).//O06730
- F-NT2RP2000027//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3) (FRAGMENT).//0.95:41: 39/MACACA FASCICULARIS (CRAB EATING MACAQUE) (CYNOMOLGUS MONKEY).//P50665
  - F-NT2RP2000032//BAX PROTEIN, CYTOPLASMIC ISOFORM GAMMA.//1.0:35:34//HOMO SAPIENS (HU-

MAN) //Q07815

F-NT2RP2000040//BASIC PROLINE-RICH PEPTIDE IB-1.//0.0024:58:36//HOMO SAPIENS (HUMAN).//P04281 F-NT2RP2000045//DNAJ PROTEIN.//1.1e-12:42:66//THERMUS AQUATICUS (SUBSP. THERMOPHILUS).//

F-NT2RP2000054//GONADOLIBERIN III PRECURSOR (GONADOTROPIN-RELEASING HORMONE III) (GN-RH-III) (LH-RH III) (LULIBERIN III).//0.20:46:36//ONCORHYNCHUS MASOU (CHERRY SALMON) (MASU SALM-ON)./P30973

F-NT2RP2000056//PROTEIN-TYROSINE PHOSPHATASE EPSILON PRECURSOR (EC 3.1.3.48) (R-PTP-EP-SILON).//1.3e-18:45:100//MUS MUSCULUS (MOUSE).//P49446

F-NT2RP2000067//HOMEOBOX PROTEIN HOX-A5 (S12-B) (FRAGMENT).//0.71:44:40//SALMO SALAR (AT-LANTIC SALMON).//P09637

F-NT2RP2000070//INSULIN.//0.94:30:43//HYSTRIX CRISTATA (CRESTED PORCUPINE).//P01328

F-NT2RP2000076//ETS-LIKE PROTEIN POINTED P1 (D-ETS-2).//0.0013:76:40//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P51022

F-NT2RP2000077//U1 SMALL NUCLEAR RIBONUCLEOPROTEIN C (U1-C).//0.24:49:40//HOMO SAPIENS (HUMAN).//P09234

F-NT2RP2000079//PLATELET FACTOR 4 (PF-4).//0.15:52:30//SUS SCROFA (PIG).//P30034

F-NT2RP2000088//HYPOTHETICAL 13.6 KD PROTEIN IN SPT4-ROM1 INTERGENIC REGION.//1.0:36:44// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53245

20 F-NT2RP2000091//HYPOTHETICAL PROTEIN HI0149 PRECURSOR.//0.22:38:47//HAEMOPHILUS INFLUEN-ZAE.//P43953

F-NT2RP2000097/VIRUS ATTACHMENT PROTEIN (O61R).//0.75:33:36//AFRICAN SWINE FEVER VIRUS (STRAIN BA71V) (ASFV).//P32510

F-NT2RP2000098

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25 F-NT2RP2000108//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.4e-09:50:70//HOMO SAPIENS (HUMAN).// P39195

F-NT2RP2000114//WISKOTT-ALDRICH SYNDROME PROTEIN (WASP).//0.024:52:44//HOMO SAPIENS (HU-MAN).//P42768

F-NT2RP2000120//5.8 KD PROTEIN IN HMC OPERON (ORF 4).//0.67:37:32//DESULFOVIBRIO VULGARIS (STRAIN HILDENBOROUGH).//P33391

F-NT2RP2000126//CHROMODOMAIN-HELICASE-DNA-BINDING PROTEIN 1 (CHD-1).//1.5e-23:94:47//HOMO SAPIENS (HUMAN).//O14646

F-NT2RP2000133//SPLICEOSOME ASSOCIATED PROTEIN 49 (SAP 49) (SF3B53).//5.6e-10:82:39//HOMO SAPIENS (HUMAN).//O15427

35 F-NT2RP2000147//CLATHRIN COAT ASSEMBLY PROTEIN AP47 (CLATHRIN COAT ASSOCIATED PROTEIN AP47) (GOLGI ADAPTOR AP-1 47 KD PROTEIN) (Ha1 47 KD SUBUNIT) (CLATHRIN ASSEMBLY PROTEIN ASSEMBLY PROTEIN COMPLEX 1 MEDIUM CHAIN)//6.7-e.89.96/98/MUS MUSCULUS (MOUSE).//P35585 F-NT2RP2000153//PEPTIDYLPROLYL ISOMERASE CYP-1 (EC 5 2 .1.8) (PEPTIDYLPROLYL CIS-TRANS ISOMERASE) (CYCLOPHILIN) (PPIASE)/1.7-e.65.136/3/JRBUGIG MALAYI //G27450

40 F-NT2RP2000157//MLO2 PROTEIN.//2.7e-06:62:40//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).// 009329

F-NT2RP2000161/IDIS3 PROTEIN HOMOLOG./I2.7e-33:173:45//CAENORHABDITIS ELEGANS./IQ17632 F-NT2RP2000173//HYPOTHETICAL 10.5 KD PROTEIN IN SODA-COMGA INTERGENIC REGION./I0.99:62:25// PACIL IUS SUPTILIS //P54499

45 F-NT2RP2000175//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.19:41:43//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//001643

F-NT2RP2000183//DIHYDROPYRIMIDINASE RELATED PROTEIN-2 (DRP-2) (NEURAL SPECIFIC PROTEIN NSP60).//4.1e-19:114:44//BOS TAURUS (BOVINE).//O02675

F-NT2RP2000195//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.99:30:33//MICROTUS PENNSYLVANI-CUS (MEADOW VOLE).//P24949

F-NT2RP2000205/MERCURIC TRANSPORT PROTEIN PERIPLASMIC COMPONENT PRECURSOR (PERI-PLASMIC MERCURY ION BINDING PROTEIN) (MERCURY SCAVENGER PROTEIN).//0.098:88:25//SH-EWANELLA PUTREFACIENS (PSEUDOMOAS PUTREFACIENS).//054463

F-NT2RP2000208//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.020:19:57//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01645

F-NT2RP2000224//PUTATIVE CUTICLE COLLAGEN C09G5.4.//0.0058:159:32//CAENORHABDITIS ELE-GANS //O09455

F-NT2RP2000232//P55-C-FOS PROTO-ONCOGENE PROTEIN (FRAGMENT).//1.0:44:38//OVIS ARIES

- (SHEEP) //O02761
- F-NT2RP2000233//GASTRIN/CHOLECYSTOKININ TYPE B RECEPTOR (CCK-B RECEPTOR) (CCK-BR).//
  0.34:53:43//CANIS FAMILIARIS (DOG).//P30552
- F-NT2RP2000239//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT).//0.019: 69:33//RATTUS NORVEGICUS (RAT).//P10164
  - F-NT2RP2000248//OVOMUCOID (FRAGMENT).//0.88:18:55//POLYPLECTRON EMPHANUM (PALAWAN PEA-COCK-PHEASANT).//P52250
  - F-NT2RP2000257//PUTATIVE MITOCHONDRIAL CARRIER YIL006W.//6.4e-09:83:37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40556
- F-NT2RP2000258//MYOSIN II HEAVY CHAIN, NON MUSCLE.//0.081:217:28//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD),//P08799
  - F-NT2RP2000270//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.4e-17:80:57//HOMO SAPIENS (HUMAN).// P39188
- F-NT2RP2000274//HYPOTHETICAL 5.8 KD PROTEIN://0.082:22:45//CLOVER YELLOW MOSAIC VIRUS (CYMV).//P16485
  - F-NT2RP2000283//HYPOTHETICAL 83.6 KD PROTEIN R05D3.2 IN CHROMOSOME III.//0.39:38:34// CAENORHABDITIS ELEGANS.//P34535
    - F-NT2RP2000288

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- F-NT2RP2000289//HYPOTHETICAL 9.4 KD PROTEIN IN RNPA-THDF INTERGENIC REGION.//0.40:38:42//ES-CHERICHIA COLL//P22847
- F-NT2RP2000297//ZINC FINGER PROTEIN 85 (ZINC FINGER PROTEIN HPF4) (HTF1).//2.3e-62:206:47//HO-MO SAPIENS (HUMAN).//Q03923
  - F-NT2RP2000298//CUTICLE COLLAGEN 12 PRECURSOR.//0.55:81:40//CAENORHABDITIS ELEGANS.// P20630
- 25 F-NT2RP2000310//RUBREDOXIN (RD).//0.13:43:41//TREPONEMA PALLIDUM.//O83956
  - F-NT2RP2000327//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:46:30//GADUS MORHUA (ATLANTIC COD).//P15996
  - F-NT2RP2000328//HYPOTHETICAL 86.6 KD PROTEIN IN PFK1-TDS4 INTERGENIC REGION.//2.0e-21:198: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53313
- 30 F-NT2RP2000329//GTP:AMP PHOSPHOTRANSFERASE MITOCHONDRIAL (EC 2.7.4.10) (AK3).//1.8e-91:155: 92//BOS TAURUS (BOVINE).//P08760
  - F-NT2RP2000337//PROTEIN A54.//0.75.48:35//VACCINIA VIRUS (STRAIN WR), AND VACCINIA VIRUS (STRAIN COPENHAGEN).//P21072
  - F-NT2RP2000346//MYELOID DIFFERENTIATION PRIMARY RESPONSE PROTEIN MYD116.//9.7e-13:114:42// MUS MUSCULUS (MOUSE).//P17564
    - F-NT2RP2000369//CALTRIN (CALCIUM TRANSPORT INHIBITOR).//0.98:47:34//MUS MUSCULUS (MOUSE).// Q09098
    - F-NT2RP2000412//SHORT NEUROTOXIN D PRECURSOR.//0.66:57:36//AIPYSURUS LAEVIS (OLIVE SEA SNAKE).//P19960
- 40 F-NT2RP2000414//HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN F (HNRNP F).//1.0e-27:96:67//HO-MO SAPIENS (HUMAN).//P52597
  - F-NT2RP2000420//ZINC FINGER PROTEIN 191.//0.16:47:38//HOMO SAPIENS (HUMAN).//O14754
- F-NTZRP2000422//PUTATIVE PHOSPHOACETYLGLUCOSAMINE MUTASE (EC 5.4.2.3) (ACETYLGLU-COSAMINE PHOSPHOMUTASE) (N-ACETYLGLUCOSAMINE-PHOSPHATE MUTASE).//3.6e-19:148:36// 45 SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//QO9887
- F-NT2RP2000438/TUBULIN GAMMA CHAIN./0.86:190.27/IRETICULOMYXA FILOSA.//PS4405 F-NT2RP2000448/VXSYSTEAU.EINDINO PROTEINI/3.7a-613:140-24/IDMOD SAPIENS (HUMAN).//P22059 F-NT2RP2000458//NEURONAL PROTEIN 3.1 (P311 PROTEIN)//1.0.46:35/HOMO SAPIENS (HUMAN).// 018612
- F-NT2RP2000498//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//0.062:25:68//HOMO SAPIENS (HUMAN).// P39194
  - F-NT2RP2000503
  - F-NT2RP2000510//TOXIN IV-5.//1.0:51:33//TITYUS BAHIENSIS (BRAZILIAN SCORPION).//P56608
  - F-NT2RP2000516//SLYX PROTEIN.//1.0:52:32//ESCHERICHIA COLI.//P30857
- 55 F-NT2RP2000523/IPHORBOLIN I (FRAGMENTS).//1.4e-06:36:47/I/HOMO-SAPIENS (HUMAN)./IP31941 F-NT2RP2000603//ALPHA/BETA-GLIADIN PRECURSOR (PROLAMIN) (CLASS A-III)./I/0.93:119:26/ITRITICUM AESTIVUM (WHEAT)./IP04723
  - F-NT2RP2000617//SPERM PROTAMINE P1 (CYSTEINE-RICH PROTAMINE).//0.056:16:62//OVIS ARIES

- (SHEEP), AND CAPRA HIRCUS (GOAT) //P04102
- F-NT2RP2000634//NEDD-4 PROTEIN (EC 6.3.2.-) (KIAA0093) (FRAGMENT).//1.8e-05:128:28//HOMO SAPI-ENS (HUMAN).//P46934
- F-NT2RP2000644//HYPOTHETICAL PROTEIN HI1566 PRECURSOR.//0.85:48:39//HAEMOPHILUS INFLUEN-7AF //P44257
- F-NT2RP2000656//EARLY GROWTH RESPONSE PROTEIN 1 (EGR-1) (NERVE GROWTH FACTOR-INDUCED PROTEIN A) (NGFI-A).//1.0:111:24//RATTUS NORVEGICUS (RAT).//P08154
  - F-NT2RP2000658//URONATE ISOMERASE (EC 5.3.1.12) (GLUCÚRONATE ISOMERASE) (URONIC ISOMERASE), //0.49:79:31//ESCHERICHIA COLI.//P42607
- 10 F-NT2RP2000668/MEROZOITE SURFACE ANTIGEN 2 PRECURSOR (MSA-2) (45 KD MEROZOITE SURFACE ANTIGEN),//0.020:115:30//PLASMODIUM FALCIPARUM (ISOLATE 3D7),//P50498
  - F-NT2RP2000678//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00085:38:68//HOMO SAPIENS (HUMAN).// P39188
  - F-NT2RP2000704//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.2e-17:55:74//HOMO SAPIENS (HUMAN).//
  - F-NT2RP2000710//ASPARTYL-TRNA SYNTHETASE (EC 6.1.1.12) (ASPARTATE-TRNA LIGASE) (ASPRS).// 8.9e-47:106:59//TREPONEMA PALLIDUM.//O83950
    - F-NT2RP2000715

- F-NT2RP2000731//CONIDIATION-SPECIFIC PROTEIN 10.//0.094:31:41//NEUROSPORA CRASSA.//P10713
- F-NT2RP2000758//!!! ALU SUBFAMILY J WARNING ENTRY !!!!/0.00027:31:74/HOMO SAPIENS (HUMAN).// P39188 F-NT2RP2000764/NIFS PROTEIN //2.7e-27:175:47//ANABAENA SP. (STRAIN PCC. 7120).//P12623
- F-NT2RP2000809//HYPOTHETICAL PROTEIN MG381 HOMOLOG.//0.91:85:25//MYCOPLASMA PNEUMONI-
  - F-NT2RP2000809//HYPOTHETICAL PROTEIN MG381 HOMOLOG.//0.91:85:25//MYCOPLASMA PNEUMONI AE.//P75219
- 25 F-NT2RP2000812//DILUTE MYOSIN HEAVY CHAIN, NON-MUSCLE (MYOSIN 5A).//2.8e-07:133:31//MUS MUS-CULUS (MOUSE).//Q99104
  - F-NT2RP2000814/40S RIBOSOMAL PROTEIN S27A.//0.93:44:38/L/COPERSICON ESCULENTUM (TOMA-TO), AND SOLANUM TUBEROSUM (POTATO).//P27083 F-NT2RP2000816//HYPOTHETICAL 88.4 KD PROTEIN B0464.7 IN CHROMOSOME III.//3.3e-21:123:39//
- OCANORHABDITIS ELEGANS.//Q03565
  F-NT2RP200819//TROPOMYOSIN 5, CYTOSKELETAL TYPE.//1.0:71:30//MUS MUSCULUS (MOUSE).//
  - P21107
  - F-NT2RP2000841//GUANINE NUCLEOTIDE RELEASING PROTEIN (GNRP).//0.0011:133:26//MUS MUSCU-LUS (MOUSE).//P27671
- 35 F-NT2RP2000842//LYSOPHOSPHATIDIC ACID RECEPTOR (EDG-2).//6.4e-13:22:95//HOMO SAPIENS (HU-MAN).//Q92633
  - F-NT2RP2000845//BOWMAN-BIRK TYPE PROTEINASE INHIBITOR (MSTI).//0.92:24:41//MEDICAGO SCUTELLATA (SNAIL MEDIC).//P80321
- F-NT2RP2000863//N-MYC PROTO-ONCOGENE PROTEIN.//0.010:148:27//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P24793
  - F-NT2RP2000880//PROBABLE TRANSLATION INITIATION FACTOR IF-2.//4.0e-100:199:94//HOMO SAPIENS (HUMAN).//O60841
  - F-NT2RP2000892//PROCOLLAGEN ALPHA 1(II) CHAIN PRECURSOR [CONTAINS: CHONDROCALCIN].//
    0.43:45:44//MUS MUSCULUS (MOUSE).//P28481
- 45 F-NT2RP2000931//MATRIN 3.//2.8e-46:104:92//RATTUS NORVEGICUS (RAT).//P43244
  - F-NT2RP2000932//2-5A-DEPENDENT RIBONUCLEASE (EC 3.1.26.-) (2-5A-DEPENDENT RNAASE) (RNASE L) (RIBONUCLEASE 4) (FRAGMENT),//3.9e-07:113:31//MUS MUSCULUS (MOUSE),//Q05921
  - F-NT2RP2000938//VOLTAGE-GATED POTASSIUM CHANNEL PROTEIN KV3.3 (KSHIIID).//0.026:59:45//RAT-TUS NORVEGICUS (RAT).//001956
- 50 F-NT2RP2000943//HYPOTHETICAL PROTEIN KIAA0079 (HA3543).//5.9e-18:161:42//HOMO SAPIENS (HU-MAN).//P53992
  - F-NT2RP2000965//INNER CENTROMERE PROTEIN (INCENP).//0.062:156:25//GALLUS GALLUS (CHICK-ENL///D53352
- F-NT2RP2000970//EC PROTEIN HOMOLOG.//1.0:50:30//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//
  - F-NT2RP2000985//HYPOTHETICAL 96.8 KD PROTEIN IN SIS2-MTD1 INTERGENIC REGION.//2.5e-06:53:47// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36159
    - F-NT2RP2000987//INSECT TOXIN 4 (INSECT TOXIN AAH IT4).//1.0:32:34//ANDROCTONUS AUSTRALIS HEC-

TOR (SAHARA SCORPION) //P21150

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- F-NT2RP2001036//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.2e-33:65:81//HOMO SAPIENS (HUMAN).//
- F-NT2RP2001044//HIRUSTASIN.//0.97:15:66//HIRUDO MEDICINALIS (MEDICINAL LEECH).//P80302
- F-NT2RP2001056//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!//1.0e-24:85:65//HOMO SAPIENS (HUMAN).//
  - F-NT2RP2001065//BOWMAN-BIRK TYPE SEED TRYPSIN AND CHYMOTRYPSIN INHIBITOR (BTCI).//0.41:50: 32//VIGNA UNGUICULATA (COWPEA).//P17734
  - F-NT2RP2001070//PROBABLE PYRIDOXAMINE 5'-PHOSPHATE OXIDASE (EC 1.4.3.5) (PNP/PMP OXIDASE)
- 10 (FPRA PROTEIN).//6.2e-18:64:48//MYXOCOCCUS XANTHUS.//P21159 F-NT2RP2001081//SYNAPTOTAGMIN IV.//7.8e-16:94:46//RATTUS NORVEGICUS (RAT).//P50232
  - F-NT2RP2001094//METALLOTHIONEIN-I (MT-I)./I/1.0:24:33//RATTUS NORVEGICUS (RAT).//P02803 F-NT2RP2001119//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/I7.5e-11:61:63//HOMO SAPIENS (HUMAN).// P30196
- 15 F-NT2RP2001127/IXE169 PROTEIN (SMCX PROTEIN) (FRAGMENTS).//1.0e-47:155:58//MUS MUSCULUS (MOUSE).//P41230
  - F-NT2RP2001137//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.10:68:39//BOS TAURUS (BOVINE).//
    - F-NT2RP2001149//!!!! ALU SUBFAMILY J WARNING ENTRY!!!!//1.1e-13:81:59//HOMO SAPIENS (HUMAN).//
- F-NT2RP2001168//PROTEIN KINASE C SUBSTRATE 80 KD PROTEIN (FRAGMENTS).//0.0071:77:33//RATTUS NORVEGICUS (RAT).//P20468
  - F-NT2RP2001173//CYTOSKELETON-ASSOCIATED PROTEIN CKAPI (TUBULIN FOLDING COFACTOR B).//
    1.0:36:41//HOMO SAPIENS (HUMAN).//Q99426
- 25 F-NT2RP2001174//ZINC FINGER PROTEIN 137.//7.2e-11:65:43//HOMO SAPIENS (HUMAN).//P52743
  - F-NT2RP2001196//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2 (EC 1.6.5.3).//1.0:95:26//CAPRA HIR-CUS (GOAT).//Q36346
  - F-NT2RP2001218//HYPOTHETICAL 59.2 KD PROTEIN IN MOB1-SGA1 INTERGENIC REGION.//0.00024:80: 23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40486
- F-NT2RP2001226//RABPHILIN-3A (FRAGMENT)//A 66-05-121-39//MUS MUSCULUS (MOUSE).//P47708 F-NT2RP2001233//ZINC FINGER PROTEIN ZFP-36 (FRAGMENT).//3-2e-61:153:56//HOMO SAPIENS (HU-MAN).//P16415
  - F-NT2RP2001245//SYNAPTONEMAL COMPLEX PROTEIN 1 (SCP-1 PROTEIN).//4.9e-05:230:21//HOMO SA-PIENS (HUMAN).//Q15431
- 35 F-NT2RP2001268//HOMEOBOX PROTEIN CEH-32.//0.23:159:25//CAENORHABDITIS ELEGANS.//Q23175 F-NT2RP2001277
  - F-NT2RP2001290/BETA-SOLUBLE NSF ATTACHMENT PROTEIN (SNAP-BETA) (SNAP-ALPHA HOMOLOG) (BRAIN PROTEIN (417) (FRAGMENT).//1.08-86:131:97//MUS MUSCULUS (MOUSE).//P28663 F-NT2RP2001295
- 40 F-NT2RP2001312//N-ACETYLGLUCOSAMINE-6-SULFATASE PRECURSOR (EC 3.1.6.14) (G6S) (GLU-COSAMINE-6-SULFATASE).//0.64:80:33//CAPRA HIRCUS (GOAT).//P50426
  - F-NT2RP2001327//TUMOR NECROSIS FACTOR, ALPHA-INDUCED PROTEIN 1, ENDOTHELIAL (B12 PRO-TEIN).//1.0e-36:118:65//HOMO SAPIENS (HUMAN).//Q13829
- F-NT2RP2001328//PROBABLE E5 PROTEIN.//1.0:46:41//HUMAN PAPILLOMAVIRUS TYPE 33.//P06426
  45 F-NT2RP2001347//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//4.5e-19:66:62//HOMO SAPIENS (HUMAN).//
  P30103
  - F-NT2RP2001366//SPERM-SPECIFIC PROTEIN PHI-1.//0.66:55:32//MYTILUS EDULIS (BLUE MUSSEL).//
    O04621
    - F-NT2RP2001378//VOLTAGE-GATED POTASSIUM CHANNEL PROTEIN KV3.3 (KSHIIID) (FRAGMENT).// 0.060:78:33//HOMO SAPIENS (HUMAN).//Q14003
  - F-NT2RP2001381//26S PROTEASE REGULATORY SUBUNIT 8 (SUG1 HOMOLOG) (XSUG1).//1.0:167:26// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P46470
    - F-NT2RP2001392//KERATIN, HIGH-SULFUR MATRIX PROTEIN, IIIA3.//0.0080:82:32//OVIS ARIES (SHEEP).// P02441
- 55 F-NT2RP2001394//POLYHOMEOTIC-PROXIMAL CHROMATIN PROTEIN.//0.024:39:53//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P39769
  - F-NT2RP2001397//G2/MITOTIC-SPECIFIC CYCLIN B2.//1.4e-46:125:78//MESOCRICETUS AURATUS (GOLD-EN HAMSTER).//P37883

- F-NT2RP2001420//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONES CP3, CP4 AND CP5) [CONTAINS: BASIC PEPTIDE IB-6; PEPTIDE P-H].//0.00018:113:38//HOMO SAPIENS (HUMAN).//P04280
- F-NT2RP2001423//HYPOTHETICAL 9.4 KD PROTEIN IN GP31-CD INTERGENIC REGION (ORF A).//0.90:23: 43//BACTERIOPHAGE T4.//P17307
- F-NT2RP2001427//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.2e-11:38:68//HOMO SAPIENS (HUMAN).//
  - F-NT2RP2001436//DYNEIN LIGHT INTERMEDIATE CHAIN 2, CYTOSOLIC (LIC53/55) (LIC-2).//0.25:124:28// RATTUS NORVEGICUS (RAT).//Q62698
- F-NT2RP2001440//14-3-3 PRÓTEIN GAMMA (PROTEIN KINASE C INHIBITOR PROTEIN-1) (KCIP-1).//4.8e-62:
  145:90//RATTUS NORVEGICUS (RAT).//P35214
- F-NT2RP2001445

F-NT2RP2001450

- F-NT2RP2001449//CLEAVAGE AND POLYADENYLATION SPECIFICITY FACTOR, 100 KD SUBUNIT (CPSF 100 KD SUBUNIT).//9.5e-118:226:95//BOS TAURUS (BOVINE).//Q10568
- 15 F-NT2RP2001467//SHORT NEUROTOXIN 1 (TOXIN V-II-1).//1.0:25:40//BUNGARUS FASCIATUS (BANDED KRAIT).//P10808
  - F-NT2RP2001506
    F-NT2RP2001511//HYPOTHETICAL 115.4 KD PROTEIN ZK757.3 IN CHROMOSOME III.//0.49:124:29//
    CAFNORHABDITIS ELEGANS.//P34681
- F-NT2RP2001520/VITAMIN D-DEPENDENT CALCIUM-BINDING PROTEIN, INTESTINAL (CABP) (CALBINDIN D9K), //0.035.71:33//HOMO SAPIENS (HUMAN), //P29377
- DSK);///.035/7133/HOMO SAPIENS (HUMAN);/P293/7
  F-NT2RP2001526
  F-NT2RP2001526//METALLOTHIONEIN-I (MT-1)://10:19:42//COLUMBA LIVIA (DOMESTIC PIGEON);//P15786
- F-NT2RP2001560//CUTICLE COLLAGEN 12 PRECURSOR.//0.0018:144:35//CAENORHABDITIS ELEGANS.//
  25 P20630
  F-NT2RP2001569//!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//2.7e-31:102:67//HOMO SAPIENS (HU-
  - MAN).//P39194 F-NT2RP2001576//SMP3 PROTEIN.//0.00016:75:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
- Q04174

  F-NTZRP2001581/TRANSMEMBRANE PROTEIN SEX PRECURSOR./I/0.040:46:36//HOMO SAPIENS (HU-
- MAN).//P51805
  F-NTZRP2001597//PROBABLE E4 PROTEIN.//0.00042:113:34//HUMAN PAPILLOMAVIRUS TYPE 5.//P06924
  - F-NT2RP2001601

    F-NT2RP2001613//HOMEOBOX PROTEIN SAX-1 (CHOX-3) (FRAGMENT).//0.14:59:32//GALLUS GALLUS
    - (CHICKEN),//P19801
      F-NT2RP2001628//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.056:140:33//SACCHAROMY-CES CEREVISIAE (BAKER'S
    - YEAST).//P32323

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- F-NT2RP2001634//ALPHA-CATENIN.//7.1e-12:152:35//DROSOPHILA MELANOGASTER (FRUIT FLY).//
  40 P35220
- F-NT2RP2001660//HYPOTHETICAL 80.4 KD PROTEIN IN SMC3-MRPL8 INTERGENIC REGION.//0.43:119:26// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40358
- F-NT2RP2001663//ALPHA ENOLASE (EC 4.2.1.11) (2-PHOSPHO-D-GLYCERATE HYDRO-LYASE) (NON-NEURAL ENOLASE) (NNE) (PHOSPHOPYRUVATE HYDRATASE)//1.2e-26:126:56//HOMO SAPIENS (HU-MAN)//P06733
- F-NT2RP2001675//HYPOTHETICAL 107.7 KD PROTEIN IN RPSO 5'REGION (ORF1).//0.25:148:25//CAMPY-LOBACTER.JEJI/NJ //Q46089
  - F-NT2RP2001677//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT).//0.010: 101:31//RATTUS NORVEGICUS (RAT).//P10164
  - F-NT2RP2001678//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.6e-18:83:61//HOMO SAPIENS (HUMAN).// P39188
    - F-NT2RP2001699//PROTEIN C14.//0:98:51:31//VACCINIA VIRUS (STRAIN COPENHAGEN).//P21045 F-NT2RP2001720//MEROZOITE SURFACE ANTIGEN 2 PRECURSOR (MSA-2) (ALLELIC FORM 1).//0.16:145:
- 30//PLASMODIUM FALCIPARUM (ISOLATE CAMP / MALAYSIA).//Q99317
  55 F-NT2RP2001721//MALE-SPECIFIC LETHAL-2 PROTEIN.//0.00090:48:39//DROSOPHILA MELANOGASTER
- (FRUIT FLY).//P50534 F-NT2RP2001740//ANNEXIN VII (SYNEXIN) (FRAGMENT).//0.50:43:25//BOS TAURUS (BOVINE).//P20072
  - F-NT2RP2001748//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-

- MENT).//0.77:111:28//HOMO SAPIENS (HUMAN).//P10162
- F-NT2RP2001762
- F-NT2RP2001813//PHOTOSYSTEM | REACTION CENTRE SUBUNIT VIII (PSI-I).//1.0:22:40//PICEA ABIES (NORWAY SPRUCE) (PICEA EXCELSA).//047040
- F-NT2RP2001839//SCY1 PROTEIN.//6.8e-17:204:27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  - F-NT2RP2001861//D15KZ1 PROTEIN (FRAGMENT).//0.31:56:39//MUS MUSCULUS (MOUSE).//Q61466 F-NT2RP2001869//CORNEODESMOSIN (S PROTEIN) (FRAGMENT).//0.97:78:30//SUS SCROFA (PIG).//
- 10 F-NT2RP2001876//ALLOGRAFT INFLAMMATORY FACTOR-1 (AIF-1) (IONIZED CALCIUM BINDING ADAPTER MOLECULE 1),//3.5e-36:106:66//HOMO SAPIENS (HUMAN),//P55008
  - F-NT2RP2001883//CATHEPSIN L (EC 3.4.22.15).//0.95:29:41//OVIS ARIES (SHEEP).//Q10991
  - F-NT2RP2001898//TYPE II INOSITOL-1,4,5-TRISPHOSPHATE 5-PHOSPHATASE PRECURSOR (EC 3.1.3.56) (5PTASE) (FRAGMENT),//1.6e-84:185:88//HOMO SAPIENS (HUMAN),//P32019
- 15 F-NT2RP2001900/IACTIN-LIKE PROTEIN ARP5.//1.16-17:180:34//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST)./P53946 F-NT2RP2001907/IHYPHAL WALL PROTEIN 1 (CELL ELONGATION PROTEIN 2).//0.13:108:27//CANDIDA AL-
  - BICANS (YEAST)./IP46593

    F-NT2RP2001926//HYPOTHETICAL 7.6 KD PROTEIN YCF33.//0.55:57:26//CYANOPHORA PARADOXA.//
  - P48273
  - F-NT2RP2001936

GARIS://P56353

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- F-NT2RP2001943//HYPOTHETICAL 57.7 KD PROTEIN IN AIP1-CTF13 INTERGENIC REGION.//1.8e-13:208: 22//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04305
- F-NT2RP2001946//HYPOTHETICAL 13.0 KD PROTEIN IN ALGR3 3'REGION.//0.59:76:28//PSEUDOMONAS AFRIGINOSA //P21485
- F-NT2RP2001947//ZINC FINGER PROTEIN DAN (N03).//0.53:68:29//RATTUS NORVEGICUS (RAT).//Q06880 F-NT2RP2001989//CHLOROPLAST 30S RIBOSOMAL PROTEIN S18://0.0016:52:34//CHLORELLA VUL-
- F-NT2RP2001976//DILUTE MYOSIN HEAVY CHAIN, NON-MUSCLE (MYOSIN 5A).//9.5e-07:201:22//MUS MUS-CULUS (MOUSE).//Q99104
- F-NT2RP2001985//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//0.016:90:32//MUS MUSCULUS (MOUSE).//
  - P05142
    F-NT2RP2001991//SODIUM- AND CHLORIDE-DEPENDENT TRANSPORTER NTT73.//8.0e-14:47:76//RATTUS
- NORVEGICUS (RAT).//008469

  F-NT2RP2002025//NG-CAM RELATED CELL ADHESION MOLECULE PRECURSOR (NR-CAM) (BRAVO).//
  2.9e-30:21142//GALLUS GALLUS (CHICKEN).//P35331
  - F-NT2RP2002032//FLOCCULANT-ACTIVE PROTEINS MO2.1 AND MO2.2://0.23:20:40//MORINGA OLEIFERA (HORSERADISH TREE) (MORINGA PTERYGOSPERMA)://P24303
- F-NT2RP2002033//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.88:27:62//HOMO SAPIENS (HUMAN).//
- F-NT2RP2002041
  - F-NT2RP2002048/JMATING PROCESS PROTEIN MID2 (SERINE-RICH PROTEIN SMS1) (PROTEIN KINASE A INTERFERENCE PROTEIN).//1.0:85:31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36027 F-NT2RP2002047
- 45 F-NT2RP2002058//DOM34 INTERACTING PROTEIN 2.//9.4e-25:165:34//SACCHAROMYCES CEREVISIAE //BAKER'S YEAST\//012220
  - F-NT2RP2002066//TIGHT JUNCTION PROTEIN ZO-1 (TIGHT JUNCTION PROTEIN 1).//5.7e-12:108:41//HOMO SAPIENS (HUMAN).//Q07157
  - F-NT2RP2002070//CYTOCHROME C OXIDASE POLYPEPTIDE II (EC 1.9.3.1) (FRAGMENT).//0.88:28:50//AS-TERINA PECTINIFERA (STARFISH).//P11958
  - F-NT2RP2002076/TRP-ASP REPEATS CONTAINING PROTEIN RBA-2.//0.0031:124:27//CAENORHABDITIS ELEGANS.//P90916
  - F-NT2RP2002078//KERATIN, GLYCINE/TYROSINE-RICH OF HAIR.//0.82:30:40//OVIS ARIES (SHEEP).// Q02958
- 55 F-NT2RP2002079//OUTER DENSE FIBER PROTEIN //0.34:41:39//HOMO SAPIENS (HUMAN)//Q14990 F-NT2RP2002099//HETEROGENOUS NUCLEAR RIBONUCLEOPROTEIN U (HNRNP U).//5.2e-08:81:48//HOMO SAPIENS (HUMAN)//000839
  - F-NT2RP2002105//COLLAGEN 1(X) CHAIN PRECURSOR.//0.0012:100:34//BOS TAURUS (BOVINE).//P23206

- F-NT2RP2002124//EARLY GROWTH RESPONSE PROTEIN 1 (EGR-1) (KROX24) (TRANSCRIPTION FACTOR ETR103) (ZINC FINGER PROTEIN 225) (AT225).//0.74:72:31//HOMO SAPIENS (HUMAN).//P18146
- F-NT2RP2002137//NEUROTOXIN B-II.//1.0:27:44//CEREBRATULUS LACTEUS (MILKY RIBBON WORM).// P01526
- 5 F-NT2RP2002154I/GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PRO-TEIN) (35 KD LECTIN) (CARBOHYDRATE BINDING PROTEIN 35) (CBP 35) (LAMINI-BINDING PROTEIN) (ILECTIN L-29) (L-34 GALACTOSIDE-BINDING LECTIN) //0.0029:112:34/MUS MUSCULUS (MOUSE) //P16110 F-NT2RP2002172
  - F-NT2RP2002185//UBIQUITIN-LIKE PROTEIN DSK2.//1.8e-07:87:40//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P48510
- 10 ER'S YEAST).//P485 F-NT2RP2002192
  - F-NT2RP2002193//CUTICLE COLLAGEN 40.//0.0062:70:37//CAENORHABDITIS ELEGANS.//P34804
  - F-NT2R-2002193/PEROXISOME ASSEMBLY PROTEIN PEX10 (PEROXIN-10).//0.00011:45:40//HOMO SAPI-ENS (HUMAN)./080683
- 15 F-NT2RP2002219
  - F-NT2RP2002231//V-TYPE SODIUM ATP SYNTHASE SUBUNIT E (EC 3.6.1.34) (NA(+)-TRANSLOCATING AT-PASE SUBUNIT E).//1.0:68:32//ENTEROCOCCUS HIRAE.//P43436
    - F-NT2RP2002235//INFECTED CELL PROTEIN ICP34.5 (NEUROVIRULENCE FACTOR ICP34.5).//0.0022:66: 45//HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN CVG-2).//P37318
- 49//HERYES SIMPLEX VIRUS (TYPE 17 STRAIN CVG-2)//F97310 F-NT2RP2002252//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RPB1) (FRAG-MENT)//0.071-110:31//CRICETULUS GRISEUS (CHINESE HAMSTER)//P11414
  - F-NT2RP2002256/CYTOCHROME P450 26 (EC 1.14.-.) (RETINOIC ACID-METABOLIZING CYTOCHROME)
    (P450RAI) (RETINOIC ACID 4-HYDROXYLASE).//3.1e-31:75:84/MUS MUSCULUS (MOUSE).//O55127
- F-NT2RP2002259//L-MYC-1 PROTO-ONCOGENE PROTEIN.//1.9e-17:41:90//HOMO SAPIENS (HUMAN).//
  25 P12524
  - F-NT2RP2002270//HYPOTHETICAL 26.0 KD PROTEIN IN CYB5-LEU4 INTERGENIC REGION.//2.1e-27:164: 36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53930
    - F-NT2RP2002292//IMMEDIATE-EARLY PROTEIN RSP40.//0.018:107:23//PSEUDORABIES VIRUS (STRAIN KAPI AN) (PRV) //P24877
- 39 F-NTZRPZ002312//PHOSPHATIDATE CYTIDYLYLTRANSFERASE (EC 2.7.7.41) (CDP-DIGLYCERIDE SYN-THETASE) (CDP-DIGLYCERIDE PYN-PROPHOSPHORYLASE) (CDP-DIACYLCLYCEROL SYNTHASE) (CDS) (CTP-PHOSPHATIDATE CYTIDYLYLTRANSFERASE) (CDP-DAG SYNTHASE)//1.4e-52:174:55//HOMO SAPIENS (HUMAN)//092903
  - F-NT2RP2002316//HISTONE H1.C6/H1.C9.//1.0:40:40//TRYPANOSOMA CRUZI.//P40269
- 35 F-NT2RP2002325//PEROXISOMAL MEMBRANE PROTEIN PMP30A (PMP31) (PEROXIN 11A).//2.2e-06:145: 26//CANDIDA BOIDINII (YEAST).//Q00316
  - F-NT2RP2002333//HYPOTHETICAL 39.1 KD PROTEIN IN RNPB-SOHA INTERGENIC REGION (ORF 3).//0.30: 86:32//ESCHERICHIA COLI.//P23524
  - F-NT2RP2002373//SYNAPSINS IA AND IB.//0.080:145:31//BOS TAURUS (BOVINE).//P17599
- F-NT2RP2002385/ENV POLYPROTEIN PRECURSOR (COAT POLYPROTEIN) [CONTAINS: KNOB PROTEIN GP7: SPIKE PROTEIN P15E: R PROTEIN] //0 021:66 28//MINK CELL FOCUS-FORMING MURINE LEUKEMIA VIRUS (ISOLATE CI-3) //P03388 F-NT2RP2002348
- F-NT2RP2002408//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//0.00030:107:37//BOS TAURUS (BO-VINE).//P02453
  - F-NT2RP2002426
  - F-NT2RP2002439//CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS).//0.00032:79:32//PLASMODIUM BERGHEI (STRAIN ANKA).//P23093
    - F-NT2RP2002442//HESA PROTEIN.//6.0e-16:163:30//PLECTONEMA BORYANUM.//P46037
  - F-NT2RP2002457

- F-NT2RP2002464//HYPOTHETICAL 60.7 KD PROTEIN C56F8.17C IN CHROMOSOME I.//9.3e-18:165:32// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10264
  - F-NT2RP2002475//CYSTEINE-RICH HEART PROTEIN (HCRHP).//0.91:45:35//HOMO SAPIENS (HUMAN).// P50238
- 55 F-NT2RP2002479//ATP-BINDING CASSETTE TRANSPORTER 7 PRECURSOR (ABC TRANSPORTER 7 PRO-TEIN).//6.8e-96:186:94//HOMO SAPIENS (HUMAN).//O75027
  - F-NT2RP2002498//HYPOTHETICAL MERCURIC RESISTANCE PROTEIN MERC.//0.65:37:45//PSEU-DOMONAS AERUGINOSA.//P04139

- F-NT2RP2002503//ZINC FINGER PROTEIN 45 (BRC1744).//1.3e-31:124:59//HOMO SAPIENS (HUMAN).// O02386
- F-NT2RP2002504//NUCLEAR PORE COMPLEX PROTEIN NUP155 (NUCLEOPORIN NUP155) (155 KD NU-CLEOPORIN) (P140) //1.2e-123:240:92//RATTUS NORVEGICUS (RAT) //P37199
- 5 F-NT2RP2002520//ACIDIC PROLINE-RICH PROTEIN HP43A PRECURSOR.//0.94:83:28//MESOCRICETUS AURATUS (GOLDEN HAMSTER).//P06680
  - F-NT2RP2002537//HYPOTHETICAL 55.1 KD PROTEIN B0416.5 IN CHROMOSOME X.//4.0e-10:194:23//
    CAENORHABDITIS ELEGANS.//Q11073
    F-NT2RP2002546
- 10 F-NT2RP2002549//G2/MITOTIC-SPECIFIC CYCLIN C13-1 (A-LIKE CYCLIN) (FRAGMENT).//0.98:65:30//DAU-CUS CAROTA (CARROT).//P25010
  - F-NT2RP2002591//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2).//2.6e-19:60:61//HOMO SAPI-ENS (HUMAN).//P51523
- F-NT2RP2002595//ANNEXIN VII (SYNEXIN).//1.2e-15:121:49//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//Q92125
- F-NT2RP2002606//PROTEIN TRANSPORT PROTEIN SEC2.//0.00034:98:31//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P17065
  - F-NT2RP2002609//HYPOTHETICAL 52.0 KD PROTEIN IN CLB6-SPT6 INTERGENIC REGION.//0.00022:79:39// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53264
- 20 F-NT2RP2002618//PROTEIN ARGININE N-METHYLTRANSFERASE 1 (EC 2.1.1.-).//6.2e-37:180:44//RATTUS NORVEGICUS (RAT).//Q63009
  - F-NT2RP2002621/MADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3) (FRAGMENT).//0.98:37: 35//LEMUR CATTA (RING-TAILED LEMUR).//Q34879
- F-NT2RP2002643//INFECTED CELL PROTEIN ICP34.5 (NEUROVIRULENCE FACTOR ICP34.5).//0.042:77:32//
  HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN MGH-10).//P37319
- F-NT2RP2002672//PROTEIN Q300.//0.0018:41:43//MUS MUSCULUS (MOUSE).//Q02722
  - F-NT2RP2002701//HYPOTHETICAL 72.5 KD PROTEIN C2F7.10 IN CHRÓMOSOME I.//3.6e-17:100:42// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09701
  - F-NT2RP2002706//MMEDIATE-EARLY PROTEIN IE180.//0.00027:139:33//PSEUDORABIES VIRUS (STRAIN KAPLAN) (PRV)./P33479
- F-NTZRP2002710/SH3-BINDING PROTEIN 3BP-1.//6.9e-09-96:40//MUS MUSCULUS (MOUSE) //P55194 F-NTZRP2002727//TUBERIN (TUBEROUS SCLEROSIS 2 HOMOLOG PROTEIN).//3.6e-20:160:36//RATTUS NORVEGICUS (RXT):/498816
  - F-NT2RP2002736 F-NT2RP2002740

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- F-NT2RP2002741//RHO1 GDP-GTP EXCHANGE PROTEIN 2.//2.0e-07:178:28//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P51862
  - F-NT2RP2002750//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!/1.6e-09:43:72//HOMO SAPIENS (HU-MAN).//P39191
- 40 F-NT2RP2002752/I/OW CALCIUM RESPONSE LOCUS PROTEIN T.//0.95:33:39//YERSINIA PSEUDOTUBER-CUI OSIS //Q00932
  - F-NT2RP2002753//ENDOGLUCANASE EG-1 PRECURSOR (EC 3.2.1.4) (ENDO-1,4-BETA-GLUCANASE) (CELLULASE).//0.71:78:33//TRICHODERMA LONGIBRACHIATUM.//Q12714
- F-NT2RP2002769//50 KD SPICULE MATRIX PROTEIN PRECURSOR //0.44:76:32//STRONGYLOCENTROTUS
  45 PURPURATUS (PURPLE SEA URCHIN).//P11994
  - F-NT2RP2002778
    - F-NT2RP2002800//CRAMBIN.//0.99:20:50//CRAMBE ABYSSINICA (ABYSSINIAN CRAMBE).//P01542
    - F-NT2RP2002839//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONE CP7) [CONTAINS: BASIC PEPTIDE P-F] (FRAGMENT).//0.010:87:31//HOMO SAPIENS (HUMAN).//P02812
  - F-NT2RP2002857//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP33).//0.00018:57:45//RAT-TUS NORVEGICUS (RAT\.//P04474
    - F-NT2RP2002862//HYPOTHETICAL 27.1 KD PROTEIN UFD4-CAP1 INTERGENIC REGION.//7.2e-27:140:40// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P33201
- F-NT2RP2002880//DNA REPAIR PROTEIN RAD32.//0.83:67:28//SCHIZOSACCHAROMYCES POMBE (FIS-55 SION YEAST).//Q09683
  - F-NT2RP2002891//HOMEOBOX PROTEIN DLX-2 (DLX-5) (FRAGMENT).//0.99:70:24//RATTUS NORVEGICUS (RAT) //064204
    - F-NT2RP2002925//ALPHA-1D ADRENERGIC RECEPTOR (ALPHA 1D-ADRENOCEPTOR) (ALPHA-1A

- ADRENERGIC RECEPTOR) //0.31:48:43//HOMO SAPIENS (HUMAN) //P25100
- F-NT2RP2002928//CELL DIVISION CONTROL PROTEIN 40.//2.8e-26:142:42//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P40968
- F-NT2RP2002929//HYPOTHETICAL 46.2 KD TRP-ASP REPEATS CONTAINING PROTEIN D2013.2 IN CHRO-MOSOME II.//2.0e-31:186:35//CAENORHABDITIS ELEGANS.//Q18964
- F-NT2RP2002939//ADENYLATE CYCLASE, TYPE V (EC 4.6.1.1) (ATP PYROPHOSPHATE-LYASE) (CA(2+)-INHIBITABLE ADENYLYL CYCLASE)//0.002298:39//CANIS FAMILIARIS (DOG)//P30803
- F-NT2RP2002959//UBIQUITIN-CONJUGATING ENZYME E2-17 KD 2 (EC 6.3.2.19) (UBIQUITIN- PROTEIN LIGASE) (UBIGUITIN CARRIER PROTEIN) (E2(17)KB 2)//2.8e-11:33.81//HOMO SAPIENS (HUMAN), MUS MUSCULUS (MOUSE), RATTUS NORVEGICUS (RAT), AND XENOPUS LAEVIS (AFRICAN CLAWED FROG).// P51869

## F-NT2RP2002979

- 15 F-NTZRP2002980//30S RIBOSOMAL PROTEIN S10.//1.1e-09:98:36//MYCOPLASMA CAPRICOLUM.//P10129 F-NTZRP2002986//RING CANAL PROTEIN (KELCH PROTEIN).//1.1e-19:141:39//DROSOPHILA MELA-NOGSTER (FBI IT FLY )//OMABO.
  - F-NT2RP2002987//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.3e-07:78:47//HOMO SAPIENS (HUMAN).//
- 20 F-NT2RP2002993//DNA-DIRECTED RNA POLYMERASE I 135 KD POLYPEPTIDE (EC 2.7.7.6) (RNA POLYMERASE I SUBUNIT)//i8.0e-77.165:85//RATTUS NORVEGICUS (RRATI/JO548/8
  - F-NT2RP2003000//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.8e-19:62:64//HOMO SAPIENS (HUMAN).//
- F-NT2RP2003034/HYPOTHETICAL PROTEIN H11458./f.10.42:35//HAEMOPHILUS INFLUENZAE.//P44204 F-NT2RP2003073/!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//0.0051:16:87//HOMO SAPIENS (HUMAN).// P30180

# F-NT2RP2003099

- F-NT2RP2003108//BASIC PROLINE-RICH PEPTIDE IB-1.//0.84:47:34//HOMO SAPIENS (HUMAN).//P04281
- F-NT2RP2003121//HYPOTHETICAL 96.7 KD PROTEIN IN STE2-FRS2 INTERGENIC REGION.//9.0e-08:99:29// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43572
  - F-NT2RP2003125/TRANSCRIPTION REGULATOR PROTEIN BACH2 (BTB AND CNC HOMOLOG 2).//9.2e-08: 134:28//MUS MUSCULUS (MOUSE).//P97303

#### F-NT2RP2003129

- F-NT2RP2003137//UBIQUITIN.//3.4e-06:70:30//NEUROSPORA CRASSA.//P13117
  - F-NT2RP2003157//HYPOTHETICAL 37.0 KD PROTEIN B0495.8 IN CHROMOSOME II.//7.8e-13:84:40// CAENORHABDITIS ELEGANS.//Q09217
- F-NT2RP2003158//26S PROTEASOME REGULATORY SUBUNIT S3 (PROTEASOME SUBUNIT P58).//3.1e-65: 155:84//HOMO SAPIENS (HUMAN).//O43242
  - F-NT2RP2003161//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//0.0011:59:42//MUS MUSCULUS (MOUSE).//P05142
    - F-NT2RP2003164//ZYXIN.//0.0037:85:36//MUS MUSCULUS (MOUSE).//Q62523
- F-NT2RP2003165//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.2e-24:77:64//HOMO SAPIENS (HUMAN).//
  45 P39194
- F-NT2RP2003177//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.55:38:39//DROSOPHILA MELA-NOGASTER (FRUIT FLY) //001643
  - F-NT2RP2003194//HYPOTHETICAL 12.5 KD PROTEIN ZK637.2 IN CHROMOSOME III.//2.3e-14:87:37//
    CAENORHABDITIS ELEGANS.//P30629
- F-NT2RP2003206//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 3 (EC 1.6.5.3).//1.0:100:28//DIDELPHIS MARSUPIALIS VIRGINIANA (NORTH AMERICAN OPOSSUM).//P41306
  - F-NT2RP2003228//DNA REPLICATION LICENSING FACTOR MCM4 (CDC21 HOMOLOG) (P1-CDC21).//9.3e-82:211:81//HOMO SAPIENS (HUMAN).//P33991
- F-NTZRP2003230//SEC14 CYTOSOLIC FACTOR (PHOSPHATIDYLINOSITOL/PHOSPHATIDYLCHOLINE
  TRANSFER PROTEIN) (PI/PC TP).//1.0:51:31//CANDIDA GLABRATA (YEAST) (TORULOPSIS GLABRATA).//
  P53989
  - F-NT2RP2003237//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/5.1e-44:66:84//HOMO SAPIENS (HUMAN).// P39194

- F-NT2RP2003243//M PROTEIN, SEROTYPE 5 PRECURSOR.//0.027:204:23//STREPTOCOCCUS PYO-GENES.//PD2977
- F-NT2RP2003265//BP4A PROTEIN.//0.95:35:34//BRASSICA NAPUS (RAPE).//P41505
- F-NT2RP2003272//ANTER-SPECIFIC PROLINE-RICH PROTEIN APG (PROTEIN CEX) (FRAGMENT).//5.5e-06:78:35//BRASSICA NAPUS (RAPE).//P40603
- F-NT2R72003277/I/NAM7 PROTEIN (NONSENSE-MEDIATED MRNA DECAY PROTEIN 1) (UP-FRAMESHIFT SUPPRESSOR 1)//1.9e-19:145-43//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)///P30771 F-NT2RP2003290
- F-NT2RP2003286//RNA 3'-TERMINAL PHOSPHATE CYCLASE (EC 6.5.1.4) (RNA-3'-PHOSPHATE CYCLASE)
- (RNA CYCLASE) //2.1e-32-137-42//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) // IQ08096 F-NT2RP2003293/ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1) // 17.7e-12:175:33//HOMO SAPIENS (HUMAN) // P51522
- F-NT2RP2003295//PTB-ASSOCIATED SPLICING FACTOR (PSF).//0.055:44:45//HOMO SAPIENS (HUMAN).// P23346
- 15 F-NT2RP2003297

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- F-NT2RP2003307/KINESIN LIGHT CHAIN (KLC).//2.0e-18:87:49//RATTUS NORVEGICUS (RAT).//P37285 F-NT2RP2003308//CROOKED NECK PROTEIN.//2.1e-91:244.67//DROSOPHILA MELANOGASTER (FRUIT FLY)./P17886
- F-NT2RP2003329//HYPOTHETICAL 54.9 KD PROTEIN C02F5.7 IN CHROMOSOME III.//5.8e-57:186:55// CAENORHABDITIS FLEGANS //P34284
- F-NT2RP2003339//SHORT NEUROTOXIN 1 (NEUROTOXIN ALPHA).//0.98:11:72//DENDROASPIS POLYLEPIS POLYLEPIS (BLACK MAMBA).//P01416
  - F-NT2RP2003347//60S RIBOSOMAL PROTEIN L38.//0.83:42:33//OSTERTAGIA OSTERTAGI.//O61570
- F-NT2RP2003367//SYNERGISTIC-TYPE VENOM PROTEIN C9S3, CHAIN 1.//1.0:37:35//DENDROASPIS AN-25 GUSTICEPS (EASTERN GREEN MAMBA).//P01408
  - F-NT2RP2003391//MRNA TRANSPORT REGULATOR MTR10.//3.3e-11:229:24//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//Q99189
  - F-NT2RP2003393//PROTOCHLOROPHYLLIDE REDUCTASE CHLB SUBUNIT (EC 1.3.1.33) (NADPH-PROTO-CHLOROPHYLLIDE OXIDOREDUCTASE CHLB SUBUNIT) (FRAGMENT).//0.94:29:34//ARAUCARIA HETERO-
- 30 PHYLLA.//P37843
  - F-NT2RP2003394
    - F-NT2RP2003401//60 KD CHAPERONIN (PROTEIN CPN60) (GROEL PROTEIN).//0.95:125:28//THERMUS AQUATICUS (SUBSP. THERMOPHILUS).//P45746
  - F-NT2RP2003433//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//9.8e-78:178:84//RATTUS NORVEGICUS (RAT).//P38378
- F-NT2RP2003445
  - F-NT2RP2003446//HYPOTHETICAL PROTEIN E-115.//0.00030:106:33//HUMAN ADENOVIRUS TYPE 2.// P03290
- F-NT2RP2003456//PHOTOSYSTEM II REACTION CENTRE M PROTEIN.//1.0:27:51//MARCHANTIA POLY-MORPHA (LIVERWORT) //P12168
- F-NTZRP2003466//LINOLEOYL-COA DESATURASE (EC 1.14.99.25) (DELTA(6)-DESATURASE).//6.7e-06:108: 32//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//008871
  - F-NT2RP2003480//TRANSCRIPTION FACTOR BF-2 (BRAIN FACTOR 2) (BF2) (CBF-2) (T-14-6).//7.2e-15:38: 50//GALLUS (GALLUS (CHICKEN).//Q98937
- 45 F-NT2RP2003499//5E5 ANTIGEN.//0.090:114:32//RATTUS NORVEGICUS (RAT).//Q63003
  - F-NT2RP2003506//NADPH-CYTOCHROME P450 REDUCTASE (EC 1.6.2.4) (CPR).//2.0e-11:91:43//SUS SCROFA (PIG).//P04175
  - F-NT2RP2003511//PARAMYOSIN, SHORT FORM (MIMIPARAMYOSIN).//0.0020:108:25//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P35416
- F-NT2RP2003513//PTB-ASSOCIATED SPLICING FACTOR (PSF).//1.2e-05:96:36//HOMO SAPIENS (HU-MAN) //P23246
  - F-NT2RP2003517//HYPOTHETICAL 12.9 KD PROTEIN CY49.27.//0.0059:22:31//MYCOBACTERIUM TUBER-CULOSIS.//Q10696
- F-NT2RP2003522//HYPOTHETICAL 10.0 KD PROTEIN.//1.0:65:30//THERMOPROTEUS TENAX VIRUS 1
  (STRAIN KRA1) (TTV1) //P19283
  - F-NT2RP2003533//LINÉ-1 REVERSE TRANSCRIPTASE HOMOLOG.//8.7e-18:94:54//HOMO SAPIENS (HU-MAN).//P08547
    - F-NT2RP2003543//SYNAPSINS IA AND IB.//0.045:101:35//RATTUS NORVEGICUS (RAT).//P09951

- F-NT2RP2003559/I/TBA2 PROTEIN (DXS9879E)//0.98:37:37/I/HOMO SAPIENS (HUMAN)//0.14657 F-NT2RP2003564//52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS-A)) (RO(SS-A)).//6.4e-35:176-44/I/HOMO SAPIENS (HUMAN)./IP19474
- F-NT2RP2003567//HYPOTHETICAL 11.2 KD PROTEIN T18D3.7 IN CHROMOSOME X.//0.72:82:34//
- F-NT2RP2003581//HOMEOBOX PROTEIN OTX1.//0.90:61:37//MUS MUSCULUS (MOUSE).//P80205 F-NT2RP2003596//ELONGATION FACTOR P (EF-P).//0.83:61:32//MYCOPLASMA GENITALIUM.//P47272
- F-NT2RP2003604//ALPHA-CATENIN./1.5e-11:152:33//DROSOPHILA MELANOGASTER (FRUIT FLY).// P35220

  F-NT2RP2003629//PHOSPHOLIPASE A2 ALPHA (EC 3.1.1.4) (PHOSPHATIDYLCHOLINE 2-ACYLHYDROLA-
- SE),/i0.97:85:27/I/CROTALUS ADAMANTEUS (EASTERN DIAMONDBACK RATTLESNAKE),//P00623
  F-NT2RP2003643/IACVI.NEURAMINATE CYTIDYI/LITRANSFERASE (EC 2.7.7.43) (CMP-N- ACETYLNEURAMINIC ACID SYNTHETASE) (CMP-NEUNAC SYNTHETASE) (CMP-SIALIC ACID SYNTHETASE),//3.9e12:94:40/NJEISSERIA MENIOSITIDIS //O57285
- 15 F-NT2RP2003668//!!!! ALU-SUBFAMILY SX WARNING ENTRY !!!!//5.0e-33:74:81//HOMO SAPIENS (HUMAN).// P39195
  - F-NT2RP2003687//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.7e-05:40:67//HOMO SAPIENS (HUMAN).//
- F-NT2RP2003691//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.5e-37:56:67//HOMO SAPIENS (HUMAN).//
  20 P39194
- F-NT2RP2003702//HYPOTHETICAL OXIDOREDUCTASE IN INLA 5'REGION (EC 1.-.-) (ORFA).//1.3e-07:98: 37/LISTERIA MONOCYTOGENES.//P25145
  - F-NT2RP2003704//GAMMA-GLUTAMYLTRANSPEPTIDASE 5 PRECURSOR (EC 2.3.2.2) (GAMMA-GLUTAMYLTRANSFERASE 5) (GGT-REL),//0.66:23:52//HOMO SAPIENS (HUMAN),//P36269
- 25 F-NT2RP2003706//GLUTAMYL AMINOPEPTIDASE (EC 3.4.11.7) (EAP) (AMINOPEPTIDASE A) (APA) (DIFFER-ENTIATION ANTIGEN OP160) //1.2e-22:197:35//HOMO SAPIENS (HUMAN) //007075
  ENTERPROPAGE // INFORMATION OF A PROPERTY OF A
  - F-NT2RP2003713/UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 6 (EC 3.1.2.15) (UBIQUITIN THIOLESTE-RASE 6) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 6) (DEUBIQUITINATING ENZYME 6) (PROTO-ON-COCENE TRE-2) I/27-6-06:119:34/I/HOMO SAPIENS (HUMAN) /I/733125
- 30 F-NT2RP2003714//ZINC FINGER PROTEIN 91 (ZINC FINGER PROTEIN HTF10) (HPF7).//6.7e-27:68:75//HO-MO SAPIENS (HUMAN).//Q05481
  - F-NT2RP2003727//HYPOTHETICAL PROTEIN MG007 HOMOLOG.//0.64:110:30//MYCOPLASMA PNEUMONI-AF //P75105
- 5-NTZRP2003737//UBIGUITIN-CONJUGATING ENZYME E2-17 KD 2 (EC 6.3.2.19) (UBIGUITIN- PROTEIN LIGASE) (UBIGUITIN CARRIER PROTEIN) (22(7)18/B) 2//11-2e-72(47:90/HOMO SAPIENS (HUMAN), MUS MUSCULUS (MOUSE), RATTUS NORVEGICUS (RAT), AND XENOPUS LAEVIS (AFRICAN CLAWED FROG).// P51689
  - F-NT2RP2003751//EXTRACELLULAR GLOBIN PRECURSOR.//0.67:68:30//PSEUDOTERRANOVA DECIPIENS (COD WORM).//P26914
- 40 F-NT2RP2003760//COATOMER GAMMA SUBUNIT (GAMMA-COAT PROTEIN) (GAMMA-COP) //1.0e-98:235: 82//BOS TAURUS (BOVINE).//P53620
  - F-NT2RP2003764//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//0.011:69:34//ORGYIA PSEUDOTSUGATA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//010341
- F-NT2RP2003769//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:33:36//EQUUS CABALLUS (HORSE).//P48663
  - F-NT2RP2003770//PHOSPHATE REGULON SENSOR PROTEIN PHOR (EC 2.7.3.-) (FRAGMENT).//0.029:35: 42//PSEUDOMONAS AERUGINOSA.//P23621
    - F-NT2RP2003777/HYPOTHETICAL 82 KD AVIRULENCE PROTEIN IN AVRBS3 REGION.//0.041:67:34//XAN-THOMONAS CAMPESTRIS (PV. VESICATORIA).//P14728
  - F-NT2RP2003781//HYPOTHETICAL 36.7 KD PROTEIN AH6.2 IN CHROMOSOME II.//4.7e-54:204:47// CAENORHABDITIS ELEGANS.//Q09201
    - F-NT2RP2003793//PSEUDO-HEVEIN (MINOR HEVEIN).//0.61:30:36//HEVEA BRASILIENSIS (PARA RUBBER TREE).//P80359
- F-NT2RP2003825//ENDOTHELIN-1 PRECURSOR (ET-1) (FRAGMENT).//1.0:35:37//CANIS FAMILIARIS (DOG).//P13206

- F-NT2RP2003840//HYPOTHETICAL 48.1 KD PROTEIN B0403.2 IN CHROMOSOME X.//2.5e-05:80:38// CAENORHABDITIS ELEGANS.//Q11076
- F-NT2RP2003857//BACTERIOCIN MICROCIN B17 PRECURSOR (MCB17).//0.54:28:50//ESCHERICHIA CO-

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F-NT2RP2003859//DROSOCIN PRECURSOR.//1.0:37:35//DROSOPHILA MELANOGASTER (FRUIT FLY).// P3a193

# F-NT2RP2003871

- F-NT2RP2003885//CUTICLE PROTEIN 32 (LM-32) (LM-ACP 32) (FRAGMENT).//1.0:28:50//LOCUSTA MIGRA-TORIA (MIGRATORY LOCUST).//P11736
  - F-NT2RP2003912//SERINE/THREONINE-PROTEIN KINASE NEK1 (EC 2.7.1.-) (NIMA-RELATED PROTEIN KINASE 1),//4.8e-110:268:80//MUS MUSCULUS (MOUSE),//P51954
- F-NT2RP2003952//AMINOPEPTIDASE B (EC 3.4.11.6) (ARGINYL AMINOPEPTIDASE) (ARGININE AMI-10 NOPEPTIDASE) (CYTOSOL AMINOPEPTIDASE IV) (AP-B).//0.00024:92:31//RATTUS NORVEGICUS (RAT).// 0.09175
  - F-NT2RP2003968//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//9.2e-05:101:36//
    XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437
- F-NT2RP2003976//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.7e-21:62:62//HOMO SAPIENS (HUMAN).//
  P39188
- F-NT2RP2003981/I/VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS8.//2.7e-08:165:22//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39702
  - F-NT2RP2003984//UNC-87 PROTEIN.//0.75:71:28//CAENORHABDITIS ELEGANS.//P37806
  - F-NT2RP2003986//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//5.3e-19:47:70//HOMO SAPIENS (HUMAN).// P39193
  - F-NT2RP2003988//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.2e-18:80:58//HOMO SAPIENS (HUMAN).//
    - F-NT2RP2004013//TRANSCRIPTION FACTOR BTF3 (RNA POLYMERASE B TRANSCRIPTION FACTOR 3).//
      1.0e-52:141:77//HOMO SAPIENS (HUMAN).//P20290
- F-NT2RP2004014//MACROPHAGE INFLAMMATORY PROTEIN-2-ALPHA (MIP2-ALPHA) (CINC-2-ALPHA).// 0.99:45:26//RATTUS NORVEGICUS (RAT).//Q10746
  - F-NT2RP2004041//SYNAPSINS IA AND IB //0.0022:51:37/IB0S TAURUS (BOVINE)./P17599
    F-NT2RP2004042//CRUSTACEAN HYPERGLYCEMIC HORMONE PRECURSOR (CHH) (FRAGMENT).//1.0:49: 28//PENAEUS VANNAMEI (PENOEID SHRIMP) (EUROPEAN WHITE SHRIMP)/0.026181
- 39 F-NT2RP2004068/ICALDESMON (CDM) //2 96-05:175:21/ICALLUS GALLUS (CHICKEN) //P12857 F-NT2RP2004081/ICADMIUM-METALLOTHIONEIN (CD-MT) ///0.93:59:23/IHELIX POMATIA (ROMAN SNAIL) (EDIBLE SNAIL) //P33187
  - F-NT2RP2004098//RAS SUPPRESSOR PROTEIN 1 (RSU-1) (RSP-1 PROTEIN) (RSP-1).//4.6e-09:121:30//HO-MO SAPIENS (HUMAN).//O15404
- 35 F-NT2RP2004124/NONHISTONE CHROMOSOMAL PROTEIN HMG-17.//0.068:63:31//GALLUS GALLUS (CHICKEN).//P02314
  - F-NT2RP2004142//HYPOTHETICAL 59.1 KD PROTEIN IN VPS15-YMC2 INTERGENIC REGION.//7.9e-05:94: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38262
  - F-NT2RP2004152//LAMIN L(I).//0.25:167:19//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P09010
- 40 F-NT2RP2004165//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.0014:124: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214
  - F-NT2RP2004170//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.012:125:30//MUS MUSCULUS (MOUSE).//P05143
- F-NT2RP2004172//HYPOTHETICAL 105.7 KD PROTEIN IN TPK3-PIR1 INTERGENIC REGION.//4.1e-26:214: 35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36051
  - F-NT2RP2004187//ZINC FINGER PROTEIN 174.//3 7e-12:76-47//HOMO SAPIENS (HUMAN).//015697 F-NT2RP2004194//HYPOTHETICAL 10.5 KD PROTEIN C31A2.13C IN CHROMOSOME 1.//0.0013:92:23// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//009730
- F-NT2RP2004196//METALLOTHIONEIN 10-II (MT-10-II),//0.92:36:36//MYTILUS EDULIS (BLUE MUSSEL),//
  PR0247
  - F-NT2RP2004207//MALE ACCESSORY GLAND SECRETORY PROTEIN 355A PRECURSOR.//0.92:62:35// DROSOPHILA SIMULANS (FRUIT FLY).//P33737
  - F-NT2RP2004226//66 KD STRESS PROTEIN (P66).//0.030:113:26//PHYSARUM POLYCEPHALUM (SLIME MOLD).//P90587
- 55 F-NT2RP2004232//PROTEIN KINASE C, MU TYPE (EC 2.7.1.-) (NPKC-MU).//2.0e-48:211:51//HOMO SAPIENS (HUMAN).//Q15139
  - F-NT2RP2004239//GLUTENIN, HIGH MOLECULAR WEIGHT SUBUNIT PW212 PRECURSOR.//0.00038:111: 36//TRITICUM AESTIVUM (WHEAT).//P08489

- F-NT2RP2004240//METALLOTHIONEIN-II (MT-II) (METALLOTHIONEIN-LIKE PROTEIN) (MT-CE).//1.0:39:28// CAENORHABDITIS ELEGANS.//P17512
- F-NT2RP2004242//RAS-RELATED PROTEIN RGP1 (GTP-BINDING REGULATORY PROTEIN RGP1).//0.0036: 64:28//ORYZA SATIVA (RICE).//P25766
- F-NT2RP2004245//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:21:42//PONGO PYGMAEUS PYG-MAEUS (BORNEAN ORANGUTAN).//P92896
  - F-NT2RP2004270//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT).//0.00023:118:33//NEPHILA CLA-VIPES (ORB SPIDER).//P46804
  - F-NT2RP2004300//PROBABLE E4 PROTEIN.//0.18:77:40//HUMAN PAPILLOMAVIRUS TYPE 8.//P06425
- F-NT2RP2004316 F-NT2RP2004321//HYPOTHETICAL 10.8 KD PROTEIN SSR2439.//1.0:50:28//SYNECHOCYSTIS SP. (STRAIN PCC 6803)//001904
  - F-NT2RP2004339//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//5.0e-33:84:77//HOMO SAPIENS (HUMAN).// P39195
- 15 F-NT2RP2004347//HYPOTHETICAL 40.9 KD PROTEIN F33H1.3 FROM CHROMOSOME II.//0.78:96:30// CAENORHABDITIS ELEGANS.//009556
  - F-NT2RP2004364//MINOR OUTER CAPSID PROTEIN (NS26) (NONSTRUCTURAL PROTEIN VP9).//0.059:143: 30//BOVINE ROTAVIRUS (STRAIN UK).//P04515
  - F-NT2RP2004365//EAMZP30-47 PROTEIN (FRAGMENT).//0.27:38:39//EIMERIA ACERVULINA.//P21959 F-NT2RP2004366//GLYCOPROTEIN L PRECURSOR.//0.64:71:28//MAREK'S DISEASE HERPESVIRUS
- (STRAIN GA) (MDHV).//P52510 F-NT2RP2004373//HISTIDINE-RICH GLYCOPROTEIN PRECURSOR (HISTIDINE-PROLINE RICH GLYCO-
- PROTEIN) (HPRG) (FRAGMENT)/0.59:50:04//ORYCTOLAGUS CUNICULUS (RABBIT)//028640
  F-NT2RP2004389//HYPOTHETICAL 70.7 KD PROTEIN F09G8.3 IN CHROMOSOME III.//4.0e-16:89:43//
  22 CAENORHABDITIS ELEGANS //P34388
  - F-NT2RP2004392
    F-NT2RP2004398//SINGLE-STRANDED NUCLEIC ACID-BINDING PROTEIN.//0.42:89:29//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST) //P10080
  - F-NT2RP2004399//SOMATOTROPIN PRECURSOR (GROWTH HORMONE).//1.0:72:34//MESOCRICETUS AU-
- 30 RATUS (GOLDEN HAMSTER).//P37886 F-NT2RP2004400

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- F-NT2RP2004412//SPERM PROTAMINE P1.//0.24:38:31//NOTORYCTES TYPHLOPS (MARSUPIAL MOLE).// P42143
- F-NT2RP2004425//SUPPRESSOR PROTEIN SRP40.//0.0087:197:22//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583
  - F-NT2RP2004463//ALPHA-2A ADRENERGIC RECEPTOR (ALPHA-2A ADRENOCEPTOR) (ALPHA-2AAR).// 1.3e-05:121:37//MUS MUSCULUS (MOUSE).//Q01338
  - F-NT2RP2004476//NICKEL-SENSITIVE T-TYPE CALCIUM CHANNEL ALPHA-1 SUBUNIT (RBE-II).//0.20:68: 36//RATTUS NORVEGICUS (RAT).//Q07652
- #0 F-NT2RP2004490//FOS-RELATED ANTIGEN 1.//0.94:59:33//HOMO SAPIENS (HUMAN).//P15407
  - F-NT2RP2004512//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4 (EC 1.6.5.3) (FRAGMENTS).//1.0:37: 32//PISASTER OCHRACEUS (SEA STAR).//P24998
  - F-NT2RP2004523//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.1e-15:57:71//HOMO SAPIENS (HUMAN).//
- 45 F-NT2RP2004538/KINESIN-LIKE PROTEIN KIF1A (AXONAL TRANSPORTER OF SYNAPTIC VESICLES).// 1.2e-48:121:60//HOMO SAPIENS (HUMAN).//O12756
  - F-NT2RP2004551/HYPOTHETICAL 7.6 KD PROTEIN (ORF 65)/I1.0:20:50//EUGLENA GRACILIS.//P32095 F-NT2RP2004568//PUTATIVE ATP-DEPENDENT RNA HELICASE C30D11.03.//5.2e-07:150:30//SCHIZOSAC-CHAROMYCES POMBE (FISSION YEAST)//0.09803
  - F-NT2RP2004580//!!!! ALU SUBFAMILY SC WARNING ENTRY!!!!//3.7e-37:100:78//HOMO SAPIENS (HU-MAN).//P39192
    - F-NT2RP2004587//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//8.2e-06: 150:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214
- F-NT2RP2004594//HYPOTHETICAL 45.3 KD PROTEIN C09F5.7 IN CHROMOSOME II.//0.84:105:24//
  55 CAENORHABDITIS ELEGANS.//Q09458
  - F-NT2RP2004600//MYRISTOYLATED ALANINE-RICH C-KINASE SUBSTRATE (MARCKS).//0.17:127:29//RAT-TUS NORVEGICUS (RAT).//P30009
    - F-NT2RP2004602//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-05:50:58//HOMO SAPIENS (HUMAN).//

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- F-NT2RP2004614//HYPOTHETICAL 11.6 KD PROTEIN.//1.0:68:33//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20561
- F-NT2RP2004655//GLYCINE-RICH RNA-BINDING PROTEIN 7.//7.0e-05:70:42//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//Q03250
- F-NT2RP2004664//HYPOTHETICAL 104.0 KD PROTEIN C32A11.03C IN CHROMOSOME I.//0.30:78:38// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10328 F-NT2RP2004675

F-NT2RP2004681

- 10 F-NT2RP2004689/HYPOTHETICAL 78.3 KD PROTEIN IN RAM2-ATP7 INTERGENIC REGION.//0.021:179:24// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P34243
  - F-NT2RP2004709//HYPOTHETICAL PROTEIN MJ0647.//0.90:39:43//METHANOCOCCUS JANNASCHII.// Q58063
  - F-NT2RP2004710//GAR2 PROTEIN.//0.085:60:30//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).// P41891
- F-NT2RP2004736//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.4e-15:97:49//HOMO SAPIENS (HUMAN).//
  - F-NT2RP2004743//MALE SPECIFIC SPERM PROTEIN MST87F.//0.43:24:41//DROSOPHILA MELANOGASTER (FRUIT FLY).//P08175
- 20 F-NT2RP2004767//36.4 KD PROLINE-RICH PROTEIN.//0.0051:88:27//LYCOPERSICON ESCULENTUM (TO-MATO).//Q00451
  - F-NT2RP2004768/SERINE/THREONINE-PROTEIN KINASE NRK1 (EC 2.7.1.-) (N-RICH KINASE 1).//9.0e-29: 166:43//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38692
    F-NT2RP2004775
- 25 F-NT2RP2004791//PROBABLE LEUCYL-TRNA SYNTHETASE (EC 6.1.1.4) (LEUCINE--TRNA LIGASE) (LEURS)./// 46-60:226:53//CAENORHABDITIS ELEGANS.//009996
  F-NT2RP2004799//SUCCINYL-COA LIGASE (GDP-FORMING), BETA-CHAIN PRECURSOR (EC 6.2.1.4) (SUCCINYL-COA SYNTHETASE, BETA CHAIN) (SCS-BETA).//Z 26-42:133.57//NEOCALLIMASTIX FRONTALIS (RUCINYL-COA SYNTHETASE, BETA CHAIN) (SCS-BETA).//Z 26-42:133.57//NEOCALLIMASTIX FRONTALIS (RUCINYL-COA SYNTHETASE, BETA CHAIN) (SCS-BETA).//Z 26-42:133.57//NEOCALLIMASTIX FRONTALIS (RUCINYL-COA SYNTHETASE).
- MEN FUNGUS).//P53587

  F-NT2RP2004802//HYPOTHETICAL 17.1 KD PROTEIN IN PUR5 3'REGION.//0.018:86:32//SACCHAROMYCES
- CEREVISIAE (BAKER'S YEAST).//P38898
  F-NTZRP2004816//H-0BETA-58 PROTEIN.//1.0e-68:145:93//MUS MUSCULUS (MOUSE).//P40336
  - F-NT2RP2004610/IPRBETA>56 PROTEIN://1.08-06:145:93//MUS MUSCULUS (MOUSE)://P40336 F-NT2RP2004841//DSRD PROTEIN://0.83:33:39//ARCHAEOGLOBUS FULGIDUS://P70742
  - F-NT2RP2004861//KERATIN, HIGH-SULFUR MATRIX PROTEIN, IIIA3A.//0.0072:41:39//OVIS ARIES (SHEEP).//P02443
- F-NT2RP2004897//METALLOTHIONEIN-LIKE PROTEIN 1.//0.99:41:41//CASUARINA GLAUCA (SWAMP OAK).//Q39511
  - F-NT2RP2004933//DEATH-ASSOCIATED PROTEIN KINASE 1 (EC 2.7.1.-) (DAP KINASE 1).//8.4e-34:102:67// HOMO SAPIENS (HUMAN).//P53355
- 40 F-NT2RP2004936//HIGH POTENTIAL IRON-SULFUR PROTEIN, ISOZYME 2 (HIPIP 2).//0.87:36:33//EC-TOTHIORHODOSPIRA VACUOLATA.//P38524
  - F-NT2RP2004959//STEM CELL FACTOR PRECURSOR (SCF) (MAST CELL GROWTH FACTOR) (MGF) (C-KIT LIGAND).//1.0:69:28//CANIS FAMILIARIS (DOG).//Q06220
- F-NTZRP2004961/ZINC FINGER PROTEIN 33A (ZINC FINGER PROTEIN KOX31) (KIAA0065) (HA0946) 45 (FRAGMENT)/I2.16-21:73:58/I/OMO SAPIENS (HUMAN)//006730 F-NTZRP2004962/IIII ALU SUBFAMILY SB WARNING ENTRY III/I/0 17:28:57/HOMO SAPIENS (HUMAN)//
  - P39189
    F-NT2RP2004967//HYPOTHETICAL 7.3 KD PROTEIN.//0.76:41:31//THERMOPROTEUS TENAX VIRUS 1
  - (STRAIN KRA1) (TTV1).//P19301
- 50 F-NT2RP2004978//SPERMATID-SPECIFIC PROTEIN T2 [CONTAINS: SPERM PROTAMINE SP2].//0.44:40:45// SEPIA OFFICINALIS (COMMON CUTTLEFISH).//P80002 E-NT2P20004987
  - F-NT2RP2004985//HYPOTHETICAL PROTEIN KIAA0144.//1.2e-51:204:57//HOMO SAPIENS (HUMAN).// Q14157
- 55 F-NT2RP2004999/LONG NEUROTOXIN 1 (ALPHA-BUNGAROTOXIN) (BGTX).//0.23:73:26//BUNGARUS MUL-TICINCTUS (MANY-BANDED KRAIT).//P01378
  - F-NT2RP2005000//ATPASE STABILIZING FACTOR 15 KD PROTEIN.//0.12:37:32//SACCHAROMYCES CERE-VISIAE (BAKER'S YEAST).//P16965

- F-NT2RP2005001//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.90:54:31//HOMO SAPIENS (HUMAN).//P22531
- F-NT2RP2005003//DOWN REGULATORY PROTEIN OF INTERLEUKIN 2 RECEPTOR.//1.6e-30:78:56//MUS MUSCULUS (MOUSE).//P15533
- F-NT2RP2005012//NPL1 PROTEIN (SEC63 PROTEIN).//0.00024:94:35//SACCHAROMYCES CEREVISIAE
  - (BAKER'S YEAST), IP14906 F-NT2RP2005018/GAG POLYPROTEIN (CORE POLYPROTEIN) [CONTAINS: CORE PROTEINS P19, P10] (FRAGMENT), I/1.091:28/IAVIAN ENDOGENOUS ROUS-ASSOCIATED VIRUS-0 (EV-2) (AVIAN RETROVIRUS RAV-0), IP009337
- 10 F-NT2RP2005020 F-NT2RP2005022//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//4.9e-11:106:35//PODOSPORA AN-SERINA//Q00808
  - F-NT2RP2005031
    F-NT2RP2005037//ANTI-SILENCING PROTEIN 1.//2.2e-32:117:55//SACCHAROMYCES CEREVISIAE (BAK-
- 15 ER'S YEAST),//P32447 F-NT2RP2005038//DNA NUCLEOTIDYLEXOTRANSFERASE (EC 2.7.7.31) (TERMINAL ADDITION ENZYME)
  - F-N1ZR-ZUDUSIS/IDNA NUCLEOTIDYLEXT INANSFERASE (EC Z.7.7.31) (TERMINAL ADDITION ENZYME) (TERMINAL DEOXYMUCLEOTIDYLTRANSFERASE) (TERMINAL TRANSFERASE)/9.3e-28:187.40//AM-BYSTOMA MEXICANUM (AXOLOTL).//O57486
  - F-NT2RP2005108//CUTICLE COLLAGEN 2.//0.33:62:38//CAENORHABDITIS ELEGANS.//P17656
- F-NT2RP2005116//PUTATIVE EUKARYOTIC TRANSLATION INITIATION FACTOR 3 ALPHA SUBUNIT (EIF-3 ALPHA).//4.0e-54:161:63//CAENORHABDITIS ELEGANS.//P34466
  - F-NT2RP2005126//CHLOROPLAST 50S RIBOSOMAL PROTEIN L27 (FRAGMENT).//0.23:46:39//PLEURO-CHRYSIS HAPTONEMOFERA.//P41552
- F-NT2RP2005139//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.016:43:37//BOS TAURUS (BOVINE).//
  P25508
- F-NT2RP2005140//HYPOTHETICAL 7.4 KD PROTEIN YCF33.//0.96:51:39//GUILLARDIA THETA (CRYPTO-MONAS PHI).//O78517
  - F-NT2RP2005144//TUBBY PROTEIN.//5.6e-08:66:45//MUS MUSCULUS (MOUSE).//P50586

- TIANA TABACUM (COMMON TOBACCO), AND SPINACIA OLERACEA (SPINACH).//P12164 F-NT2RP2005162//HYPOTHETICAL 54.2 KD PROTEIN IN ERP5-ORC6 INTERGENIC REGION.//1.2e-33:139: 51//SACCHAROMYCES CEREVISIAE (BAKERS Y BASTI//P3882)
  - F-NT2RP2005168//HETEROGENOUS NUCLEAR RIBONUCLEOPROTEIN U (HNRNP U).//2.8e-33:102:61//HO-MO SAPIENS (HUMAN).//Q00839
  - F-NT2RP2005204/IDNA DAMAGE TOLERANCE PROTEIN RHC31 (RAD31 HOMOLOG).//3.9e-28:141:42//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q066/24
- F-NT2RP2005227

  F-NT2RP2005239/TRNA SPLICING PROTEIN SPL1.//2.0e-38:117:64//CANDIDA ALBICANS (YEAST).///P87185

  F-NT2RP2005254//OMEGA-AGATOXIN IB (OMEGA-AGA-IB) (FRAGMENT).//0.26:29-48//AGELENOPSIS AP-
  - ERTA (FUNNEL-WEB SPIDER).//P15970 F-NT2RP2005270//HOMEOBOX PROTEIN HOX-A4 (CHOX-1.4).//0.037:82:34//GALLUS GALLUS (CHICKEN).//
  - P17277
    F-NT2RP2005276//LONG-CHAIN-FATTY-ACID—COA LIGASE 4 (EC 6.2.1.3) (LONG-CHAIN ACYL-COA SYN-
- 45 THETASE 4) (LACS 4).//2.0e-59:174:61//RATTUS NORVEGICUS (RAT).//035547 F-NT2RP2005287//ZINC FINGER PROTEIN 26 (ZINC FINGER PROTEIN KOX20) (FRAGMENT).//1.5e-05:27:
  - 70//HOMO SAPIENS (HUMAN).//P17031
  - F-NT2RP2005288//PROBABLE RUBREDOXIN HUPI.//1.0:42:28//RHIZOBIUM LEGUMINOSARUM (BIOVAR VI-CIAE).//P28151
- 50 F-NT2RP2005289//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.1e-21:75:70//HOMO SAPIENS (HUMAN).// P39193
  - F-NT2RP2005293//TRANSLATION INITIATION FACTOR IF-2.//0.58:170:24//HELICOBACTER PYLORI (CAMPYLOBACTER PYLORI).//P55972
- F-NT2RP2005315//CUTICLE COLLAGEN 7 (FRAGMENT).//0.091:65:38//CAENORHABDITIS ELEGANS.//
  55 P18832
- F-NT2RP2005325//CHROMOGRANIN A PRECURSOR (CGA) (PITUITARY SECRETORY PROTEIN I) (SP-I) [CONTAINS: PANCREASTATIN: WE-141.//9.5e-09:98:39//HOMO SAPIENS (HUMAN).//P10645
  - F-NT2RP2005336//HYPOTHETICAL 68.7 KD PROTEIN IN STB1-MCK1 INTERGENIC REGION.//0.00011:124:

28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P42846

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- F-NT2RP2005344//PROBABLE CALCIÚM-TRANSPORTING ATPASE 4 (EC 3.6.1.38).//4.7e-21:92:52//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q12675 F-NT2RP2005354
- F-NT2RP2005358//MYOSIN IC HEAVY CHAIN.//0.012:91:39//ACANTHAMOEBA CASTELLANII (AMOEBA).//
  - F-NT2RP2005360//ACROSIN PRECURSOR (EC 3.4.21.10).//0.0022:73:36//ORYCTOLAGUS CUNICULUS
  - (RABBIT).//P48038 F-NT2RP2005393//HYPOTHETICAL 25.9 KD PROTEIN AH6.3 IN CHROMOSOME II.//0.00085:135:28//
- CAENORHABDITIS ELEGANS //Q09202
  - F-NT2RP2005407//SQUALENE MONOOXYGENASE (EC 1.14.99.7) (SQUALENE EPOXIDASE) (SE).//0.96: 109:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32476
- F-NT2RP2005436//EXTENSIN PRECURSOR (PROLINE-RICH GLYCOPROTEIN).//0.0011:54:42//ZEA MAYS (MAIZE).//P14918
- 15 F.NT2RP2005441//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//0.039:182:29//MUS MUSCULUS (MOUSE).//P05142 F.NT2RP2005453
  - F-NT2RP2005457//NADH-UBIQUINONE OXIDOREDUCTASE SUBUNIT B14.5B (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-B14.5B) (CI-B14.5B) (I/4.0e-10:124:37//BOS TAURUS (BOVINE) //O02827
- 20 F-NT2RP2005464//HÝPOTHETICAL 9.5 KD PROTEIN.//0.96:42:33//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20553
- F-NT2RP2005465//MITOCHONDRIAL CARRIER PROTEIN RIM2.//4.6e-09:92:42//SACCHAROMYCES CERE-VISIAE (BAKER'S YEAST).//P38127
- F-NT2RP2005472//HYPOTHETICAL PROTEIN BB0129.//0.76:80:32//BORRELIA BURGDORFERI (LYME DIS-25 EASE SPIROCHETE).//051155
  - F-NT2RP2005476//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.0e-31:39:89//HOMO SAPIENS (HUMAN).//
  - F-NT2RP2005490//METALLOTHIONEIN-II (MT-II).//0.14:27:33//SCYLLA SERRATA (MUD CRAB).//P02806 F-NT2RP2005491//DNA-DIRECTED RNA POLYMERASE SUBUNIT | (EC 2.7.7.6).//0.95:45:31//METHANO-
- 30 COCCUS JANNASCHII.//Q58785 F-NT2RP2005495//HYPOTHETICAL 10.8 KD PROTEIN IN GP30-RIII INTERGENIC REGION.//0.99:68:30//BAC-
- TERIOPHAGE T4.//Q02407
  - F-NT2RP2005496//ZINC FINGER PROTEIN 135.//1.46-54:120:59//HOMO SAPIENS (HUMAN).//P52742 F-NT2RP2005498//PROTEIN PHOSPHATASE PP2A, 55 KD REGULATORY SUBUNIT, ALPHA ISOFORM (PRO-
- TEIN PHOSPHATASE PPZA B SUBUNIT ALPHA ISOFORM) (ALPHA-PR55),//9.5e-76:146:86/RATTUS NOR-VEGICUS (RAT),//P36876 F-NTZRP2005501/GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PRO-
- TEIN) (35 KD LECTIN) (CARBOHYDRATE BINDING PROTEIN 35) (CBP 35) (LAMININ-BINDING PROTEIN) (LECTIN L-29) (L-31) (GALACTOSIDE-BINDING PROTEIN) (GALBP),//0.025/70-40/HOMO SAPIENS (HU-MAN),//P17931
- F-NT2RP2005509//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//1.0:166:27//GALLUS GALLUS (CHICK-FN) //P02457
  - F-NT2RP2005520//CHROMOSOME ASSEMBLY PROTEIN XCAP-E.//7.9e-45:118:79//XENOPUS LAEVIS (AF-RICAN CLAWED FROG).//P50533
- 45 F-NT2RP2005525//50S RIBOSOMAL PROTEIN L11.//1.0:47:27//BORRELIA BURGDORFERI (LYME DISEASE SPIROCHETE).//051354
  - F-NT2RP2005531//PROTEIN-TYROSINE PHOSPHATASE MEG1 (EC 3.1.3.48) (PTPASE-MEG1) (MEG).//9.8e-13:84:45//HOMO SAPIENS (HUMAN).//P29074
  - F-NT2RP2005539//RING CANAL PROTEIN (KELCH PROTEIN).//4.9e-10:90:33//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
  - F-NTZRP2005540/INUCLEOTIDE BINDING PROTEIN EXPZ/I0.36:119:21/IBACILLUS SUBTILIS.//P98115 F-NTZRP2005549/IHYPOTHETICAL 32.0 KD PROTEIN C18C10.10 IN CHROMOSOME III.//6.0e-39:179:46// CAENDRHABDITIS ELEGANS.//Q09253
- F-NT2RP2005557//HYPOTHETICAL 23.7 KD PROTEIN C13G6.14 IN CHROMOSOME I.//4.9e-06:90:35// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09790
  - F-NT2RP2005581
    F-NT2RP2005600//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.014;37;40//HOMO SAPIENS (HUMAN).//

#### P02811

F-NT2RP2005605//GONADOLIBERIN I PRECURSOR (LHRH I) (LUTEINIZING HORMONE RELEASING HORMONE I) (GONADOTROPIN RELEASING HORMONE I) (GNRH I) (LULIBERIN I) (FRAGMENT).//0.64:26:42// MACACA MILATTA (RHESUS MACAQUE).//P55247

F-NT2RP2005620//HYPOTHETICAL 45.1 KD PROTEIN IN RPS5-ZMS1 INTERGENIC REGION.//8.7e-31:138: 49//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47160

F-NT2RP2005622//NEUROTOXIN-LIKE PROTEIN STR1 (ANATOXIN AAH STR1).//0.39:22:40//ANDROCTONUS AUSTRALIS HECTOR (SAHARA SCORPION).//P80950

F-NT2RP2005635//HYPOTHETICAL 80.7 KD PROTEIN IN ERG7-NMD2 INTERGENIC REGION.//5.8e-43:144: 56//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38795

F-NT2RP2005637/VPU PROTEIN (U ORF PROTEIN).//0.91:33:45//CHIMPANZEE IMMUNODEFICIENCY VI-RUS (SIV/CPZ)) (CIV).//P17286

F-NT2RP2005640//METALLOTHIONEIN-LIKE PROTEIN LSC54.//0.63:41:31//BRASSICA NAPUS (RAPE).// P43402

## 15 F-NT2RP2005645

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F-NT2RP2005651//OCTAMER-BINDING TRANSCRIPTION FACTOR 3A (OCT-3A) (OCT-4).//0.0023:50:42//HO-MO\_SAPIENS (HIJMAN) //Q01860

F-NT2RP2005654//HYPOTHETICAL 48.6 KD PROTEIN IN BET1-PAN1 INTERGENIC REGION.//6.1e-16:76:44// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P40564

F-NT2RP2005669//METALLOTHIONEIN-II (MT-II).//0.76:16:50//SCYLLA SERRATA (MUD CRAB).//P02806 F-NT2RP2005675//PUTATIVE ORAL CANCER SUPPRESSOR (DELETED IN ORAL CANCER-1).//6.5e-26:116: 54//MESOCRICETUS AURATUS (GOI DEN HAMSTER).//P49119

54/MESOCRICE IDS AURATUS (GOLDEN HAMSTER)/P49119
F-NT2RP2005683//HYPOTHETICAL PROTEIN HI0275.//0.17:50:40//HAEMOPHILUS INFLUENZAE.//P43975
F-NT2RP2005690//PYRROLINE-5-CARBOXYLATE REDUCTASE (EC 1.5.1.2) (P5CR) (P5C REDUCTASE).//

1.3e-16:75:30//PISUM SATIVUM (GARDEN PEA).//Q04708 F-NT2RP2005694//HYPOTHETICAL PROTEIN KIAA0032.//9.6e-11:135:34//HOMO SAPIENS (HUMAN).//

Q15034 F-NT2RP2005701//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE M) [CONTAINS: PEPTIDE P-D] (FRAG-MENT.//O.084:158:32//HOMO SAPIENS (HUMAN).//P10161

F-NT2RP2005712//METALLOTHIONEIN-II (MT-II).//0.19:14:50//STENELLA COERULEOALBA (STRIPED DOL-PHIN).//P14425

F-NT2RP2005719//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENT).//1.0:36:41//ORYCTOLAGUS CUNICULUS (RABBIT).//P02456

F-NT2RP2005722//ZINC FINGER PROTEIN ZFP-36 (FRAGMENT).//7.8e-37:131:62//HOMO SAPIENS (HU-MAN).//P16415

F-NT2RP2005723//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//0.98:23:60//HOMO SAPIENS (HUMAN).//

F-NT2RP2005726//HYPOTHETICAL PROTEIN TP0375.//0.98:30:43//TREPONEMA PALLIDUM.//083390 F-NT2RP2005732//PERIOD CLOCK PROTEIN (FRAGMENT).//0.41:20:55//DROSOPHILA ROBUSTA (FRUIT

FLY)/I/Q03296 F-NT2RP2005741//SMR1 PROTEIN PRECURSOR (VCS-ALPHA 1).//0.38:58:36//RATTUS NORVEGICUS (RAT).//P13432

F-NT2RP2005748//ZINC FINGER PROTEIN KOX23 (FRAGMENT).//0.026:19:68//HOMO SAPIENS (HUMAN).//

45 F-NT2RP2005752//PROCOLLAGEN ALPHA 1(III) CHAIN PRECURSOR.//0.90:101:31//HOMO SAPIENS (HU-MAN).//P02461

F-NT2RP2005753//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.50:22:59//HOMO SAPIENS (HUMAN).//P30808

F-NT2RP2005763//PUTATIVE ATP-DEPENDENT RNA HELICASE STE13.//4.7e-14:108:37//SCHIZOSACCHA-ROMYCES POMBE (FISSION YEAST).//Q09181

F-NT2RP2005767//NONHISTONE CHROMOSOMAL PROTEIN 6B.//4.1e-08:65:32//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST).//P11633

F-NT2RP2005773//PYRROLINE-5-CARBOXYLATE REDUCTASE (EC 1.5.1.2) (P5CR) (P5C REDUCTASE).//
1.2e-14:65:61//HOMO SAPIENS (HUMAN).//P32322

55 F-NTZRP2005775/NEUROLYSIN PRECURSOR (EC. 3.4.24.16) (NEUROTENSIN ENDOPEPTIDASE) (MITO-CHONDRIAL OLIGOPEPTIDASE M) (MICROSOMAL ENDOPEPTIDASE) (MEP).//1.3e-103:199.90//ORYCTOLAGUS CUNICULUS (RABBIT)./P42875

F-NT2RP2005781//SALIVARY ACIDIC PROLINE-RICH PHOSPHOPROTEIN 1/2 PRECURSOR (PRP-1 / PRP-

- 3) (PRP-2 / PRP-4) (PIF-F / PIF-S) (PROTEIN A / PROTEIN C) [CONTAINS: PEPTIDE P-C]. //0.090:73:36//HOMO SAPIENS (HUMAN). //P02810
- F-NT2RP2005784/TRANS-ACTING TRANSCRIPTIONAL PROTEIN ICPO (IMMEDIATE-EARLY PROTEIN IET10) (VMW110) (ALPHA-0 PROTEIN).//3.5e-06:79:37//HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN 17).// P0R393
- F-NT2RP2005804//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//1.8e-07:43:55//OWENIA FUSI-FORMIS //P21260
  - F-NT2RP2005812//HYPOTHETICAL 39.3 KD PROTEIN IN GCN4-WBP1 INTERGENIC REGION.//6.3e-14:143: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40004
- F-NTZRP2005815//FERROCHELATASE (EC.4.99.1.1) (PROTOHEME FERRO-LYASE) (HEME SYNTHETASE).// 0.0017:123:37//MYCOBACTERIUM AVIUM.//007401
  - F-NT2RP2005835//SHP1 PROTEIN.//1.2e-08:135:26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P34223
- F-NT2RP2005841//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//0.23:28:53//HOMO SAPIENS (HUMAN).//P22532
  - F-NT2RP2005853//HYPOTHETICAL 8.5 KD PROTEIN IN ASIA-MOTA INTERGENIC REGION.//0.99:33:48// BACTERIOPHAGE 14.//P22917
    - F-NT2RP2005857//CHROMOSOME ASSEMBLY PROTEIN XCAP-C.//8.6e-84:235:66//XENOPUS LAEVIS (AF-RICAN CLAWED FROG).//P50532
- 20 F-NT2RP2005859//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.017:60:40//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643
  - F-NT2RP2005868//ATP SYNTHASE B' CHAIN PRECURSOR (EC 3.6.1.34) (SUBUNIT II).//0.28:121:28//SPINA-CIA OLERACEA (SPINACH).//P31853
- F-NT2RP2005886//MICRONUCLEAR LINKER HISTONE POLYPROTEIN (MIC LH) [CONTAINS: LINKER HIS-TONE PROTEINS ALPHA, BETA, DELTA AND GAMMA]///0.80:130:28//TETRAHYMENA THERMOPHILA// PA0631
  - F-NT2RP2005890

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- F-NT2RP2005901//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.35:18:44//DROSOPHILA YAKUBA (FRUIT FLY).//P03933
- 30 F-NT2RP2005908//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.0e-28:61:65//HOMO SAPIENS (HUMAN).// P39194
  - F-NT2RP2005933//PERIOD CLOCK PROTEIN (P230) (FRAGMENT).//1.7e-11:85:49//ACETABULARIA MEDI-TERRANEA (MERMAID'S WINE GLASS).//P12347
    - F-NT2RP2005942//POLY(A) POLYMERASE (EC 2.7.7.19) (PAP) (POLYNUCLEOTIDE ADENYLYLTRANS-FERASE).//7.2e-59:216:58//BOS TAURUS (BOVINE).//P25500
      - F-NT2RP2005980//HYPOTHETICAL 11.5 KD PROTEIN IN RSP8A-AST1 INTERGENIC REGION.//1.0:49:34// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38185
      - F-NT2RP2006023//DNA REPAIR PROTEIN RECN (RECOMBINATION PROTEIN N) (FRAGMENT).//1.0:40:45//
        VIBRIO CHOLERAE.//P52118
- 40 F-NT2RP2006038//HYPOTHETICAL 30.2 KD PROTEIN C02F5.4 IN CHROMOSOME III.//4.0e-11:90:34// CAENORHABDITIS ELEGANS.//P34281
  - F-NT2RP2006043/LAMININ BETA-1 CHAIN VARIANT (LAMININ BETA-1-2 CHAIN) (FRAGMENT).//0.00067:73: 38//GALLUS GALLUS (CHICKEN).//Q01636
- F-NT2RP2006052//METALLOTHIONEIN-I (MT-I).//0.19:31:38//CERCOPITHECUS AETHIOPS (GREEN MON-KEY) (GRIVET).//P02797
  - F-NT2RP2006069//COLLAGEN ALPHA 2(I) CHAIN (FRAGMENTS).//1.0:66:34//RATTUS NORVEGICUS (RAT).// P02466
    - F-NT2RP2006071//RESTIN.//0.40:156:29//GALLUS GALLUS (CHICKEN).//O42184
  - F-NT2RP2006098//HYPOTHETICAL 21.7 KD PROTEIN IN TUP1-ABP1 INTERGENIC REGION.//0.99:95:20// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P25651
  - F-NT2RP2006100//LONG NEUROTOXIN 4 (ALPHA-NEUROTOXIN).//0.94:43:34//OPHIOPHAGUS HANNAH (KING COBRA) (NAJA HANNAH).//P80156
    - F-NT2RP2006103//50S RIBOSOMAL PROTEIN L32.//0.40:36:38//SYNECHOCYSTIS SP. (STRAIN PCC 6803).// P73014
- 55 F-NT2RP2006106//CUTICLE COLLAGEN 1.//0.28:85:29//CAENORHABDITIS ELEGANS.//P08124
  - F-NT2RP2006141/HYPOTHETICAL 72.5 KD PROTEIN C2F7.10 IN CHROMOSOME I.//1.9e-08:57:42// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09701 F-NT2RP2006186

- F-NT2RP2006184//HYPOTHETICAL 11.2 KD PROTEIN IN CSGC-MDOG INTERGENIC REGION PRECURSOR //0.95.87;26//FSCHERICHIA.COL //P75917
- F-NT2RP2006186//MICROTUBULE-ASSOCIATED PROTEIN 2.//0.088:124:33//MUS MUSCULUS (MOUSE).//
- F-NT2RP2006196//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/4.0e-05:49:61//HOMO SAPIENS (HUMAN).//
  - F-NT2RP2006200//PROCOLLAGEN ALPHA 2(V) CHAIN PRECURSOR://0.0013:205:32//HOMO SAPIENS (HU-MAN) //P05997
- F-NT2RP2006219//GONADAL PROTEIN GDL.//3.5e-18:158:37//DROSOPHILA MELANOGASTER (FRUIT FLY)//P22468
  - F-17.1/F22406
    F-NT2RP2006237//FIBRINOGEN- AND IG-BINDING PROTEIN PRECURSOR (MRP PROTEIN).//0.79:103:28//
    STREPTOCOCCUS PYOGENES.//P30141
  - F-NT2RP2006238//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//4.7e-07:127:39//MUS MUSCULUS (MOUSE).//P05143
- 15 F-NT2RP2006258//PROBABLE E5 PROTEIN://0.78:47:34//RHESUS PAPILLOMAVIRUS TYPE 1 (RHPV 1).// P24834
  - F-NT2RP2006261//PENAEIDIN-3A PRECURSOR (P3-A).//0.61:35:40//PENAEUS VANNAMEI (PENOEID SHRIMP) (EUROPEAN WHITE SHRIMP).//P81058
- F-NT2RP2006275//ELECTROMOTOR NEURON-ASSOCIATED PROTEIN 2 (FRAGMENT).//1.2e-28:59:57//
  TORPEDO CALIFORNICA (PACIFIC ELECTRIC RAY).//P14401
  - F-NT2RP2006312//HIGH-MOBILITY-GROUP PROTEIN (NONHISTONE CHROMOSOMAL PROTEIN).//1.6e-06: 53:35//TETRAHYMENA PYRIFORMIS.//P40625
    - F-NT2RP2006320//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN) (FRAGMENT).//0.90:24:41//
      HUMAN IMMUNODEFICIENCY VIRUS TYPE 1 (BH5 ISOLATE) (HIV-1).//P04612
  - F-NT2RP2006321///!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.0051:25:76//HOMO SAPIENS (HUMAN).// P39193
    - F-NT2RP2006323/WISKOTT-ALDRICH SYNDROME PROTEIN (WASP).//0.84:33:39//HOMO SAPIENS (HU-MAN).//P42768
    - F-NT2RP2006333/MYOTOXIN 3 PRECURSOR (CROTAMINE 3).//0.56:37:40//CROTALUS DURISSUS TER-RIFICUS (SOUTH AMERICAN RATTLESNAKE).//P24333
  - F.-NTZRP2006334/ISUCCINYL-COA LIGASE (IDP-FORMING), ALPHA-CHAIN 3 PRECURSOR (EC 6.2.1.4) (SUCCINYL-COA SYNTHETASE, ALPHA CHAIN 3)/I/O.0097-46.41/INTICHOMONAS VAGINALIS./P534011 F.-NTZRP2006366/INONSPECIFIC LIPID-TRANSFER PROTEIN 4.3 PRECURSOR (LTP 4.3)/I/O.18:75:29/IHOR-DELM VALIGARE (BARLEY) //IO24282
- F-NT2RP2006393//OMEGA-CONOTOXIN MVIIC PRECURSOR (FRAGMENT).//0.82:15:66//CONUS MAGUS (MAGUS CONE).//P37300
  - F-NT2RP2006436//ANTERIOR-RESTRICTED HOMEOBOX PROTEIN (RATHKE POUCH HOMEO BOX).//1.4e-08:50:50//MUS MUSCULUS (MOUSE).//Q61658
- F-NT2RP2006441/METALLOTHIONEIN-LIKE PROTEIN 1.//0.99:22:54//MIMULUS GUTTATUS (SPOTTED MONKEY FLOWER) (YELLOW MONKEY FLOWER) //P20238
  - F-NTZRP2006454/SPERM PROTAMINE P1.//0.60.47:36//TACHYGLOSSUS ACULEATUS ACULEATUS (AUS-TRALIAN ECHIDNA).//P35311 F-NTZRP2006456
- F-NT2RP2006464/PHOTOSYSTEM | IRON-SULFUR CENTER (PHOTOSYSTEM | SUBUNIT VII) (9 KD POLYPEPTIDE) (PSI-C) //0.91:79:30//SYNECHOCOCCUS SP. (STRAIN PCC 7002) (AGMENELLUM QUADRU-PLICATUM //P31047)
  - F-NT2RP2006467//PUTATIVE CUTICLE COLLAGEN F55C10.3.//0.15:53:35//CAENORHABDITIS ELEGANS.// Q21184
  - F-NT2RP2006472//HYPOTHETICAL 19 KD PROTEIN (ORF 167).//0.33:98:26//MARCHANTIA POLYMORPHA
- F-NT2RP2006534

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- F-NT2RP2006554//ANTI-SIGMA F FACTOR ANTAGONIST (STAGE II SPORULATION PROTEIN AA).//0.91:50: 34//BACILLUS SPHAFRICUS //032723
- F-NT2RP2006565//SECRETORY CARRIER-ASSOCIATED MEMBRANE PROTEIN 1 (SCAMP 37).//6.0e-66:93: 96//RATTUS NORVEGICUS (RAT).//P56603
- F-NT2RP2006571//CYTOCHROME P450 2B10 (EC 1.14.14.1) (CYPIB10) (TESTOSTERONE 16-ALPHA HY-DROXYLASE) (P450-16-ALPHA) (CLONE PF3/46) //4.5e-40:138:57//MUS MUSCULUS (MOUSE).//P12791
  - F-NT2RP2006573//SPERM PROTAMINE P1 (CYSTEINE-RICH PROTAMINE).//0.53:46:39//BOS TAURUS (BO-

VINE).//P02318

- F-NT2RP2006598//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.3e-12:44:77//HOMO SAPIENS (HUMAN).//
- F-NT2RP3000002//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.4e-19:60:63//HOMO SAPIENS (HUMAN).// P39192
- F-NT2RP3000031//HYPOTHETICAL 89.8 KD PROTEIN F41H10.6 IN CHROMOSOME IV.//2.1e-39:210:42// CAENORHABDITIS FLEGANS.//Q20296
- F-NT2RP3000046//POSSIBLE THIOPHENE AND FURAN OXIDATION PROTEIN THDF.//1.4e-25:149:44// PSEUDOMONAS PUTIDA.//P25755
- f-NT2RP3000047//NPL4 PROTEIN.//4.7e-48:275:38//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P33755
  - F-NT2RP3000050//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//3.2e-72:232:59//HOMO SA-PIENS (HUMAN).//P51522
- F-NT2RP3000055/MALE SPECIFIC SPERM PROTEIN MST84DB.//0.26:57:36//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643
  - F-NT2RP3000068//HYPOTHETICAL 182.0 KD PROTEIN IN NMD5-HOM6 INTERGENIC REGION.//0.0014:66: 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47170
    - F-NT2RP3000072//HYPOTHETICAL 6.7 KD PROTEIN IN NOHA-CSPI INTERGENIC REGION //0.95:49:30//ES-CHERICHIA COLI //P77695
- 20 F-NT2RP3000080//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.1e-17:64:68//HOMO SAPIENS (HUMAN).// P39188
  - F-NT2RP3000085//BIOTIN CARBOXYLASE (EC 6.3.4.14) (A SUBUNIT OF ACETYL-COA CARBOXYLASE (EC 6.4.1.2)) (ACC).//4.4e-43:169:51//BACILLUS SUBTILIS.//P49787
    - F-NT2RP3000092//CELL DIVISION CONTROL PROTEIN 1.//0.00016:103:31//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P40986
  - F-NT2RP3000109//ACYL CARRIER PROTEIN HOMOLOG (ACP).//0.76:83:28//MYCOPLASMA GENITALIUM.// P47529
    - F-NT2RP3000134
      F-NT2RP3000142//GAR2 PROTEIN.//0.00098:241:20//SCHIZOSACCHAROMYCES POMBE (FISSION
- 30 YEAST).//P41891 F-NT2RP3000149//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.0014:33:36//PONGO PYGMAEUS ABE-
  - LII (SUMATRAN ORANGUTAN).//P92694
    F-NTZRP3000186//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//8.3e-15:36:83//HOMO SAPIENS (HUMAN).//
    P39188
- 35 F-NT2RP3000197//HYPOTHETICAL 6.0 KD PROTEIN IN THI12 5'REGION.//0.91:21:52//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53820
  - F-NT2RP3000207/IGLUCOAMYLASE S1/S2 PRECURSOR (EC 3.2.1.3) (GLUCAN 1.4-ALPHA- GLUCOSI-DASE) (1,4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//0.026:209:27//SACCHAROMYCES CEREVISIAE (BAKERS YEAST).//908640
- 40 F-NT2RP3000220//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//1.0:26:42//HOMO SAPIENS (HUMAN).//P30808
  - F-NT2RP3000233//RING CANAL PROTEIN (KELCH PROTEIN).//2.1e-42:249:39//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- F-NT2RP3000235//HOMEOBOX PROTEIN H40 (FRAGMENT),//0.55:45:40//APIS MELLIFERA (HONEYBEE).//
  P15858
  - F-NT2RP3000247//HYPOTHETICAL PROTEIN KIAA0218.//1.7e-82:123:69//HOMO SAPIENS (HUMAN).//
    O93075
  - F-NT2RP3000251//SERINE PROTEINASE STUBBLE (EC 3.4.21.-) (STUBBLE-STUBBLOID PROTEIN) //1.0:53: 33//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q05319
- 50 F-NT2RP3000252//HYPOTHETICAL 40 KD GTP-BINDING PROTEIN IN RIBOSOMAL PROTEIN GENE CLUSTER 5/REGION.//2.2e-06:96:32//HALOBACTERIUM CUTIRUBRUM.//P17103
  - F-NT2RP3000255//HISTONE H1.1 (FRAGMENT),//0.95:71:33//BOS TAURUS (BOVINE),//P02253 F-NT2RP3000267//HYPOTHETICAL 21.1 KD PROTEIN IN SSR-SERA INTERGENIC REGION (O182),//0.38:77:
- 33//ESCHERICHIA COLI./P09160

  55 F-NT2RP3000299/MYQSINI C HEAVY CHAIN./I1.2e-11:147:34//ACANTHAMOEBA CASTELLANII (AMOEBA).//
- P10569

  F-N12RP3000299//MYOSIN IC HEAVY CHAIN.//1.2e-11:147:34//ACAN HAMOEBA CAS FELLANII (AMOEBA).//
  - F-NT2RP3000312//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.64:216: 29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214

F-NT2RP3000320/ITRANSLATION INITIATION FACTOR IF-2/I/S-2-05:184:22/IAQUIFEX AEOLICUS //067825 F-NT2RP3000324/IHYPOTHETICAL PROTEIN HI1036.//0.69:64:35/IHAEMOPHILUS INFLUENZAE.//P44097 F-NT2RP3000333/WIR14 PROTEIN.//0.35:51:41/ITRITICUM AESTIVUM WHEATI./001482

F-NT2RP3000341//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.1e-30:57:80//HOMO SAPIENS (HUMAN).//

#### E-NT2RP3000348

F-NT2RP3000350//HYPOTHETICAL 40 KD GTP-BINDING PROTEIN IN RIBOSOMAL PROTEIN GENE CLUSTER 5'REGION.//0.0011:77:35//HALOBACTERIUM CUTIRUBRUM.//P17103

F-NT2RP3000359//GTP:AMP PHOSPHOTRANSFERASE MITOCHONDRIAL (EC 2.7.4.10) (AK3).//1.2e-97:222:

84//BOS TAURUS (BOVINE).//P08760

F-NT2RP3000361//PRE-MRNA SPLICING FACTOR PRP6.//2.2e-08:128:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P19735

F-NT2RP3000366//RAS-RELATED PROTEIN RAB-18 //2.1e-107:206:99//MUS MUSCULUS (MOUSE)://P35293 F-NT2RP3000393//HOMEOBOX PROTEIN HOX-C4 (HOX-3E) (CP19).//0.0023:36:52//HOMO SAPIENS (HU-MAN)://P09017

F-NT2RP3000397//PUTATIVE PRE-MRNA SPLICING FACTOR RNA HELICASE (DEAH BOX PROTEIN 13).//
5.5e-27:116:44//MUS MUSCULUS (MOUSE).//O35286

F-NT2RP3000403//PRE-MRNA PROCESSING PROTEIN PRP40.//0.00044:67:34//SACCHAROMYCES CERE-VISIAE (BAKER'S YEAST) //P33203

F-NT2RP3000418//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49): ENDONUCLASE.I/I2 2e-16:228:34//MUS MUSCULUS (MOUSE).//P11369

F-NT2RP3000433//IIII ALU SUBFAMILY J WARNING ENTRY IIII//1.7e-17:79:55//HOMO SAPIENS (HUMAN).//

F-NT2RP3000439//HYPOTHETICAL 46.4 KD PROTEIN IN FFH-GRPE INTERGENIC REGION.//9.8e-10:201: 26//ESCHERICHIA COLL//P37908

F-NT2RP3000441//PROTEIN-EXPORT MEMBRANE PROTEIN SECG HOMOLOG.//0.91:48:35//MYCOBACTE-RIUM LEPRAE.//P38388

F-NT2RP3000449//HOMEOBOX PROTEIN HOX-B8 (CHOX-2.4) (FRAGMENT).//1.0:42:33//GALLUS GALLUS (CHICKEN).//P23681

## 30 F-NT2RP3000451

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F-NT2RP3000456//COLLAGEN ALPHA 1(I) CHAIN

(FRAGMENTS).//0.00018:178:36//RATTUS NORVEGICUS (RAT).//P02454

F-NT2RP3000484//METALLOTHIONEIN-III (MT-III) (GROWTH INHIBITORY FACTOR) (GIF).//0.098:40:27//BOS TAURUS (BOVINE).//P37359

35 F-NT2RP3000487//SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185).//0.00037:16:81/NOLVOX CART-ERI.//P21997

# F-NT2RP3000512

F-NT2RP3000526//HYPOTHETICAL NIN REGION PROTEIN ORF56.//0.51:37:43//BACTERIOPHAGE LAMB-DA.//P03769

40 F-NT2RP3000527//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//1.0e-16:234:30//HOMO SA-PIENS (HUMAN).//P51522

F-NT2RP3000531//POLIOVIRUS RECEPTOR PRECURSOR (CD155 ANTIGEN).//3.4e-15:192:30//HOMO SA-PIENS (HUMAN).//P15151

F-NT2RP3000542//CYTOCHROME C OXIDASE POLYPEPTIDE II (EC 1.9.3.1) (FRAGMENT).//0.60:51:39//AS-45 TERINA PECTINIFERA (STARFISH).//P11958

F-NT2RP3000561//HYPOTHETICAL ATP-BINDING PROTEIN MJ0423.//0.79:53:32//METHANOCOCCUS JAN-NASCHII.//057866

F-NT2RP3000562//ACCESSORY GLAND PEPTIDE PRECURSOR (PARAGONIAL PEPTIDE B).//0.99:26:34// DROSOPHILA MAURITIANA (FRUIT FLY), AND DROSOPHILA SIMULANS (FRUIT FLY).//O18666

F-NT2RP3000578/HYPOTHÉTICAL 49.8 KD PROTEIN IN RPL14B-GPA1 INTERGENIC REGION.//1.5e-26:127: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38755 E-NT2P3000582

#### F-N12RP3000582

F-NT2RP3000584//METALLOTHIONEIN-II (MT-II).//0.28:27:29//MUS MUSCULUS (MOUSE).//P02798

F-NT2RP3000590//UVS-2 PROTEIN.//4.8e-10:113:33//NEUROSPORA CRASSA.//P33288

55 F-NT2RP3000592//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII-135) (TAFII-130), (TAFII-130),

F-NT2RP3000596//YEMANUCLEIN-ALPHA.//1.8e-05:98:34//DROSOPHILA MELANOGASTER (FRUIT FLY).// P25992

F-NT2RP3000599//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//0.00095:90:37//HOMO SAPIENS (HUMAN).//Q15428

F-NT2RP3000603//5E5 ANTIGEN.//1.0e-09:181:34//RATTUS NORVEGICUS (RAT).//Q63003

F-NT2RP3000605//STEROL REGULATORY ELEMENT BINDING PROTEIN-1 (SREBP-1) (STEROL REGULA-TORY ELEMENT-BINDING TRANSCRIPTION FACTOR 1).//0.00098:76:34//HOMO SAPIENS (HUMAN).//

F-NT2RP3000622//HYPOTHETICAL PROTEIN MG096 HOMOLOG 5 (P02\_ORF427).//0.15:52:36//MYCOPLAS-MA PNFIJMONIAF //P75277

F-NT2RP3000624//HYPOTHETICAL PROTEIN KIAA0256.//5.4e-16:222:31//HOMO SAPIENS (HUMAN).//

F-NT2RP3000628

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F-NT2RP3000632//ZINC FINGER PROTEIN 90 (ZFP-90) (ZINC FINGER PROTEIN NK10).//2.0e-16:52:63//MUS MUSCILLUS (MOUSE).//061967

F-NT2RP3000644///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//6.7e-40:102:79//HOMO SAPIENS (HU-MAN).//P39194

F-NT2RP3000661//HYPOTHETICAL 139.1 KD PROTEIN C08B11.3 IN CHROMOSOME II.//6.0e-08:83:36// CAENORHABDITIS ELEGANS.//Q09441

F-NT2RP3000665/HOMEOBOX PROTEIN PROPHET OF PIT-1 (PROP-1) (PITUITARY SPECIFIC HOMEODO-MAIN FACTOR).//0.13:48:35//HOMO SAPIENS (HUMAN).//075360

20 F-NT2RP3000685//HYPOTHETICAL 33.5 KD PROTEIN IN CAT1 5'REGION (ORFY).//0.26:202:23//CLOSTRID-IUM KLUYVERI.//P38943

F-NT2RP3000690//INORGANIC PYROPHOSPHATASE (EC 3.6.1.1) (PYROPHOSPHATE PHOSPHO- HYDRO-LASE) (PPASE).//0.99:131:26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P00817

F-NT2RP3000736//HYPOTHETICAL 28.7 KD PROTEIN IN RNR3-ARC15 INTERGENIC REGION.//3.5e-27:211: 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40516

F-NT2RP3000739//HYPOTHETICAL 33.5 KD PROTEIN C1D4.02C IN CHROMOSOME I.//6.0e-23:114:42// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10149

F-NT2RP3000742/I-PHOSPHATIDYLINOSITOL-4,5-BISPHOSPHATE PHOSPHODIESTERASE DELTA 1 (EC 3.1.4.11) (PLC-DELTA-1) (PHOSPHOLIPASE C-DELTA-1) (PLC-III).//6.7e-12:85:36//RATTUS NORVEGICUS (RATT)/PHO888

F-NT2RP3000753//CELL SURFACE GLYCOPROTEIN 1 PRECURSOR (OUTER LAYER PROTEIN B) (S-LAYER PROTEIN 1).//0.00011:208:28//CLOSTRIDIUM THERMOCELLUM.//Q06852

F-NT2RP3000759/IADP-RIBOSYLATION FACTOR 6//8.1e-28:141:38//GALLUS (CHICKEN),//P26990 F-NT2RP3000815//CYTOCHROME C-551 (C551) (CYTOCHROME C8).//0.24:45:37//PSEUDOMONAS DENITRIFICANS //P001103

F-NT2RP3000825//ALPHA-LACTALBUMIN (LACTOSE SYNTHASE B PROTEIN (EC 2.4.1.22)).//0.82:51:39// MACROPUS RUFOGRISEUS (RED-NECKED WALLABY).//P07458

F-NT2RP3000826//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.025:79:37//BOS TAURUS (BOVINE).// P25508

40 F-NT2RP3000836/HYPOTHETICAL PROTEIN IN KSGA 3'REGION (ORF L5) (FRAGMENT).//0.85:36:47/MYC-OPLASMA CAPRICOLUM //P43040

F-NT2RP300841/IUDP-GLICURONOSYLTRANSFERASE 1-7 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) (UGT1-7) (UGT1-07) (UGT1-7) (UGT1-7) (UGT1-7) (UGT1-7) (UGT1-8) (MGUSE)//062452

45 F-NT2RP3000845//PUTATIVE SERINE/THREONINE-PROTEIN KINASE P78 (EC 2.7.1.-).//5.2e-72:247:61//HO-MO SAPIENS (HUMAN).//P27448

F-NT2RP300847/HYPOTHETICAL PROTEIN KIAA0161./I0.037:55:30/HOMO SAPIENS (HUMAN).//P50876
F-NT2RP3000850//IIII ALU SUBFAMILY SQ WARNING ENTRY IIII/I7.4e-31:90:75/HOMO SAPIENS (HUMAN).//
P39194

F-NT2RP3000852//HYDROPHOBIC SEED PROTEIN (HPS).//0.33:23:69//GLYCINE MAX (SOYBEAN).//P24337 F-NT2RP3000859//IMMEDIATE-EARLY PROTEIN.//3.6e-07:189:25//HERPESVIRUS SAIMIRI (STRAIN 11).// 0.01042

F-NT2RP3000865

F-NT2RP3000868//MYOSIN HEAVY CHAIN, CARDIAC MUSCLE ISOFORM (FRAGMENT).//1.4e-09:232:28// GALLUS GALLUS (CHICKEN).//P29616

F-NT2RP3000869//CUTICLE COLLAGEN 2.//4.5e-08:58:46//CAENORHABDITIS ELEGANS.//P17656

F-NT2RP3000875//HOMEOBOX PROTEIN CDX-2 (CAUDAL-TYPE HOMEOBOX PROTEIN 2).//0.90:62:37// MUS MUSCULUS (MOUSE).//P43241

- F-NT2RP3000901//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//0.99:124:33//BOS TAURUS (BOVINE).// P02453
- F-NT2RP3000904
- F-NT2RP3000917//DHP1 PROTEIN.//6.5e-60:229:55//SCHIZOSACCHAROMYCES POMBE (FISSIOI
- YEAST),//P40848 F-NT2RP3000919//HYPOTHETICAL 33.5 KD PROTEIN C1D4.02C IN CHROMOSOME I.//2.4e-19:159:34// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//O10149
  - F-NT2RP3000968//40S RIBOSOMAL PROTEIN S15A.//3.7e-48:73:98//HOMO SAPIENS (HUMAN), AND RAT-TUS NORVEGICUS (RAT).//P39027
- 10 F-NT2RP3000980//COPA/INCA PROTEIN (REPA3 PROTEIN).//0.24-19:47/IESCHERICHIA COLI.//P13946 F-NT2RP3000994//MATERNAL EFFECT PROTEIN STAUFEN.//1.46-10:78:48//DROSOPHILA MELA-NOGASTER (FRUIT FLY1///P25159
  - F-NT2RP3001004//HYPOTHETICAL 7.6 KD PROTEIN B0563.8 IN CHROMOSOME X.//0.70:50:32//
- 15 F-NT2RP3001007

F-NT2RP3001109

- F-NT2RP3001055//N-TERMINAL ACETYLTRANSFERASE COMPLEX ARD1 SUBUNIT HOMOLOG.//1.3e-05: 138:28//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P36416
- F-NT2RP3001057//ZINC FINGER PROTEIN 45 (BRC1744).//4.0e-28:141:51//HOMO SAPIENS (HUMAN).//
- F-NT2RP3001081//HYPOTHETICAL 46.4 KD PROTEIN T16H12.5 IN CHROMOSOME III.//3.8e-08:144:29// CAENORHABDITIS ELEGANS.//P34568
  - F-NT2RP3001084//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT).//3.4e-06:217:32//NEPHILA CLA-VIPES (ORB SPIDER).//P46804
- F-NT2RP3001096//SYNAPTONEMAL COMPLEX PROTEIN SC65.//1.1e-30:244:33//RATTUS NORVEGICUS (RAT).//O64375
- F-NT2RP3001107//ARYLSULFATASE F (EC 3.1.6.-) (ASF) (FRAGMENT).//0.041:47:44//HOMO SAPIENS (HU-MAN).//P54793
- F-NT2RP3001111//MALE SPECIFIC SPERM PROTEIN MST84DC.//0.17:28:39//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01644
- 30 NOGASTER (FRUIT FLY).//Q01644 F-NT2RP3001113//INVOLUCRIN.//0.00036:192:23//MUS MUSCULUS (MOUSE).//P48997
  - F-N12RP3001115///NVOLUCRIN.//0.00036:192:23///MUS MUSCULUS (MOUSE).//P4899/ F-NT2RP3001115
    - F-NT2RP3001116//AMINOPEPTIDASE G (EC 3.4.11.-) (FRAGMENT).//0.99:29:51//STREPTOMYCES LIVI-DANS.//054340
- 35 F-NT2RP3001119//COLLAGEN ALPHA 4(IV) CHAIN (FRAGMENT).//0.0015:73:39//BOS TAURUS (BOVINE).//
  - F-NT2RP3001120//ZINC FINGER PROTEIN ZFP-36 (FRAGMENT).//1.3e-57:229:52//HOMO SAPIENS (HU-MAN)//P16415
- F-NT2RP3001126//HYPOTHETICAL 91.2 KD PROTEIN IN RPS4B-SCH9 INTERGENIC REGION.//2.8e-07:83:
  49 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38888
  - F-NT2RP3001133//CALCIUM BINDING PROTEIN.//2.0e-08:171:32//DICTYOSTELIUM DISCOIDEUM (SLIME
  - F-NT2RP3001140//F-SPONDIN PRECURSOR.//2.0e-147:244:97//RATTUS NORVEGICUS (RAT).//P35446 F-NT2RP3001147//TROPOMYOSIN 2 (TMII).//0.11:159:23//SCHISTOSOMA MANSONI (BLOOD FLUKE).//
- 45 P42638 FATOPPROMATERIAL PROTEIN TO (10.0) OF 40.05 (MAIL) MARGONIA (MOLECL)
  - F-NT2RP3001150//OCTAPEPTIDE-REPEAT PROTEIN T2.//6.2e-09:163:25//MUS MUSCULUS (MOUSE).// Q06666
  - F-NT2RP3001155//DNA POLYMERASE ALPHA-BINDING PROTEIN (POB1/CTF4 PROTEIN) (CHROMOSOME REPLICATION PROTEIN CHL15).//4.1e-05:244:23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// 011454
  - F-NT2RP3001176//LEUKOSIALIN PRECURSOR (LEUCOCYTE SIALOGLYCOPROTEIN) (SIALOPHORIN) (CD43) (LY 48) (B CELL DIFFERENTIATION ANTIGEN LP-3)//0.21:136:26//MUS MUSCULUS (MOUSE).//
- F-NT2RP3001214//SAP1 PROTEIN.//0.058:133:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  55 P30055
- F-NT2RP3001216//CYLICIN I (MULTIPLE-BAND POLYPEPTIDE I) (FRAGMENT).//2.1e-08:137:33//HOMO SA-PIENS (HUMAN).//P35663
  - F-NT2RP3001221//GAMMA-BUTYROBETAINE,2-OXOGLUTARATE DIOXYGENASE (EC 1.14.11.1) (GAMMA-

- BUTYROBETAINE HYDROXYLASE).//4.2e-05:131:26//PSEUDOMONAS SP. (STRAIN AK-1).//P80193
- F-NT2RP3001232//HYPOTHETICAL PROTEIN PRECURSOR IN CS5 3'REGION (FRAGMENT).//0.75:57:31// ESCHERICHIA COLL.//P33792
- F-NT2RP3001236//TRANSFORMING PROTEIN MAF.//0.017:136:30//AVIAN MUSCULOAPONEUROTIC FIB-ROSARCOMA VIRUS AS42.//P23091
  - F-NT2RP3001239/ELECTROMOTOR NEURON-ASSOCIATED PROTEIN 1 (FRAGMENT).//4.2e-55:221.49//
    TORPEDO CALIFORNICA (PACIFIC ELECTRIC RAY).//P14400
    F-NT2RP3001745
- F-NT2RP3001253//TROPOMYOSIN 2, MUSCLE THORACIC ISOFORM (TROPOMYOSIN I).//0.0042:142:24// DROSOPHILA MELANOGASTER (FRUIT FLY).//P09491
  - F-NT2RP3001260//COLLAGEN ALPHA 4(IV) CHAIN PRECURSOR://0.0011:89:43//HOMO SAPIENS (HU-MAN).//P53420
  - F-NT2RP3001268//ZINC FINGER PROTEIN 45 (BRC1744).//9.0e-29:194:44//HOMO SAPIENS (HUMAN).// Q02386
- 15 F-NT2RP3001272/HYPOTHETICAL 75.2 KD PROTEIN C13F4.08C IN CHROMOSOME I.//8.2e-17:183:26// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O10199
  - F-NT2RP3001274//SERINE/THREONINE PROTEIN PHOSPHATASE 5 (EC 3.1.3.16) (PP5) (PROTEIN PHOS-PHATASE T) (PPT) (FRAGMENT)//1.7e-09:78:39//MUS MUSCULUS (MOUSE)//060676
    - F-NT2RP3001281//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.7e-08:38:71//HOMO SAPIENS (HUMAN).// P39188
      - F-NT2RP3001297//HYPOTHETICAL PROTEIN KIAA0281 (HA6725).//2.2e-57:159:70//HOMO SAPIENS (HU-MAN).//092556
      - F-NT2RP3001307//SPERM PROTAMINE P1.//0.21:46:39//ORNITHORHYNCHUS ANATINUS (DUCKBILL PLAT-YPUS).//P35307
  - F-NT2RP3001318 F-NT2RP3001325//ENHANCER OF RUDIMENTARY HOMOLOG.//1.0:73:24//BRACHYDANIO RERIO (ZE-BRAFISH) (ZEBRA DANIO).//1098874
    - F-NT2RP3001338/IZINC FINGER PROTEIN 29 (ZINC FINGER PROTEIN KOX26) (FRAGMENT).//0.0021:56: 35//HOMO SAPIENS (HUMAN).//P17037
- 30 F-NT2RP3001339//CITRON PROTEIN.//3.6e-06:90:33//MUS MUSCULUS (MOUSE).//P49025
  - F-NT2RP3001340//HYPOTHETICAL PROTEIN UL61.//7.2e-11:202:34//HUMAN CYTOMEGALOVIRUS (STRAIN AD169).//P16818
- F-NT2RP3001355/JTRICARBOXYLATE TRANSPORT PROTEIN PRECURSOR (CITRATE TRANSPORT PRO-TEIN) (CTP) (TRICARBOXYLATE CARRIER PROTEIN)//.76-16-12/33/JHOMO SAPIENS (HUMAN)//P53007 39 F-NT2RP3001356//RAS-RELATED PROTEIN RABA (FRAGMENT).//0.00041:66:28//DICTYOSTELIUM DISCOI-DEUM (SUIME MOLD)//P34141
  - F-NT2RP3001374

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- F-NT2RP3001383//PTB-ASSOCIATED SPLICING FACTOR (PSF).//2.5e-06:190:32//HOMO SAPIENS (HU-MAN).//P23246
- 40 F-NT2RP3001384/ICHORION PROTEIN S15.//0.00079-94:37//DROSOPHILA VIRILIS (FRUIT FLY).//P13424 F-NT2RP3001392//VPU PROTEIN (ORF-X PROTEIN) (UPX PROTEIN).//1.0.22:45//CAPRINE ARTHRITIS EN-CEPHALITIS VIRUS (CAEV).//P31834
  - F-NT2RP3001396//HYPOTHETICAL 8.1 KD PROTEIN (ORF4).//1.0:37:32//STRAWBERRY MILD YELLOW EDGE-ASSOCIATED VIRUS (SMYEAV).//Q00848
- 45 F-NT2RP3001398/KRUEPPÈL-RELATED ZINC FINGER PROTEIN 2 (HKR2 PROTEIN) (FRAGMENT).//1.9e-08:45:37/HOMO SAPIENS (HUMAN).//P10073
  - F-NT2RP3001399//SSU72 PROTEIN.//7.3e-18:84:52//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P53538
  - F-NT2RP3001407//SCY1 PROTEIN.//1.5e-08:143:25//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P53009
    - F-NT2RP3001420//HYPOTHETICAL 7.9 KD PROTEIN://0.25:41:26//VACCINIA VIRUS (STRAIN COPENHA-GEN) //P20542
    - F-NT2RP3001426//DNAJ PROTEIN.//7.5e-15:78:43//HAEMOPHILUS INFLUENZAE.//P43735
  - F-NT2RP3001427//WERNER SYNDROME HELICASE.//3.6e-13:159:33//HOMO SAPIENS (HUMAN).//Q14191 F-NT2RP3001428//NUCLEOPROTEIN TPR.//1.8e-53:117:99//HOMO SAPIENS (HUMAN).//P12270
  - F-NT2RP3001432//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 (EC 1.6.5.3) (FRAGMENT).//0.96:52: 21//TARSIUS SYRICHTA (TARSIER).//Q36151
    - F-NT2RP3001447//HYPOTHETICAL 5.5 KD PROTEIN IN REPLICATION ORIGIN REGION (ORF1).//0.96:45:35//

ESCHERICHIA COLL/P14505

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- F-NT2RP3001449//HOMEOBOX PROTEIN SAX-1 (CHOX-3) (FRAGMENT).//0.0043:53:43//GALLUS GALLUS (CHICKEN).//P19601
- F-NT2RP3001453//MALE SPECIFIC SPERM PROTEIN MST84DB://0.0048:65:40//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643
- F-NT2RP3001457//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS28.//0.55:121:20//SACCHA-ROMYCES CEREVISIAE (BAKER'S YEAST).//Q02767
  - F-NT2RP3001459//MYOSIN IC HEAVY CHAIN.//0.10:126:34//ACANTHAMOEBA CASTELLANII (AMOEBA).// P10569
- F-NT2RP3001472//NONHISTONE CHROMOSOMAL PROTEIN 6A.//3.0e-14:87:43//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST).//P11632 F-NT2RP3001490//METALLOTHIONEIN-LIKE PROTEIN LSC54.//1.0:39:35//BRASSICA NAPUS (RAPE).//
  - P43402
- F-NT2RP3001495//UBIQUITIN--PROTEIN LIGASE RSP5 (EC 6.3.2.-),//3.3e-14:148:35//SACCHAROMYCES 15 CEREVISIAE (BAKER'S YEAST),//P39940
- F-NT2RP3001497//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.13:44:38//BOS TAURUS (BOVINE).//
  - F-NT2RP3001527//SPERM PROTAMINE P1.//0.35;29;37//DIDELPHIS MARSUPIALIS VIRGINIANA (NORTH AMERICAN OPOSSUM), AND MONODELPHIS DOMESTICA (SHORT-TAILED GREY OPOSSUM), //P35305
- 20 F-NT2RP3001529//HYPOTHETICAL 43.3 KD GTP-BINDING PROTEIN IN DACB-RPMA INTERGENIC RE-GION.//3.3e-21:125:37//ESCHERICHIA COLI.//P42641
  - F-NT2RP3001538//HNF3/FH TRANSCRIPTION FACTOR GENESIS (WINGED HELIX PROTEIN CWH-3).//0.13: 53:39//GALLUS GALLUS (CHICKEN).//P79772
- F-NT2RP3001554//ELECTROMOTOR NEURON-ASSOCIATED PROTEIN 2 (FRAGMENT), 1/2, 3e-48:137:52// 25 TORPEDO CALIFORNICA (PACIFIC ELECTRIC RAY).//P14401
  - F-NT2RP3001580//GERM CELL-LESS PROTEIN.//8.2e-18:100:42//DROSOPHILA MELANOGASTER (FRUIT FLY) //O01820
- F-NT2RP3001587//UBIQUITIN-ACTIVATING ENZYME E1-LIKE (POLYMERASE-INTERACTING PROTEIN 2).// 2.0e-47:188:51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P52488
- F-NT2RP3001589//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/7.4e-41:87:80//HOMO SAPIENS (HUMAN).// P39193
  - F-NT2RP3001607//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:49:32//DICENTRARCHUS LABRAX (EUROPEAN SEA BASS),//Q36362
  - F-NT2RP3001608//EXTENSIN PRECURSOR (PROLINE-RICH GLYCOPROTEIN).//0.0013:177:25//ZEA MAYS (MAIZE).//P14918 F-NT2RP3001621//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.84:29:37//DROSOPHILA MELA-
    - NOGASTER (FRUIT FLY).//Q01645 F-NT2RP3001629//RAS-RELATED C3 BOTULINUM TOXIN SUBSTRATE 1 (P21-RAC1) (FRAGMENTS).//0.91:
    - 57:24//CAVIA PORCELLUS (GUINEA PIG).//P80236 F-NT2RP3001634//!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//8.9e-11:73:54//HOMO SAPIENS (HUMAN).//
    - F-NT2RP3001642//HYPOTHETICAL PROTEIN KIAA0210.//1.1e-12:117:29//HOMO SAPIENS (HUMAN).//
    - Q92609 F-NT2RP3001646//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//0.0092:69:34//ORGYIA PSEUDOTSUGATA
- 45 MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//O10341 F-NT2RP3001671//RING CANAL PROTEIN (KELCH PROTEIN).//0.0042:55:41//DROSOPHILA MELA-
  - NOGASTER (FRUIT FLY).//Q04652 F-NT2RP3001672
    - F-NT2RP3001676//GTP-BINDING PROTEIN LEPA (FRAGMENT).//1.2e-15:56:62//PSEUDOMONAS FLUO-RESCENS //P26843
- F-NT2RP3001678//SPIDROIN 2 (DRAGLINE SILK FIBROIN 2) (FRAGMENT),//0.054;187;31//NEPHILA CLA-VIPES (ORB SPIDER), //P46804
  - F-NT2RP3001679//HYPOTHETICAL 68.7 KD PROTEIN ZK757.1 IN CHROMOSOME III.//1.5e-07:63:44// CAENORHARDITIS ELEGANS //P34679
- 55 F-NT2RP3001688//GLUCOAMYLASE S1 PRECURSOR (EC 3.2.1.3) (GLUCAN 1.4-ALPHA-GLUCOSIDASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE) (GAI).//1.0:83:28//SACCHAROMYCES DIASTATICUS (YEAST).//P04065
  - F-NT2RP3001690//MYOSIN HEAVY CHAIN, CARDIAC MUSCLE BETA ISOFORM.//0.021:247:24//HOMO SA-

PIENS (HUMAN).//P12883

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- F-NT2RP3001698
- F-NT2RP3001708//TWISTED GASTRULATION PROTEIN PRECURSOR.//7.7e-12:73:43//DROSOPHILA MEL-ANOGASTER (FRUIT FLY).//P54356
- F-NT2RP3001712//CEC-1 PROTEIN.//1.9e-07:121:29//CAENORHABDITIS ELEGANS.//P34618
  - F-NT2RP3001716//SALIVARY GLUE PROTEIN SGS-3 PRECURSOR.//0.89:54:40//DROSOPHILA SIMULANS (FRUIT FLY).//P13729
    - F-NT2RP3001724//CHROMODOMAIN-HELICASE-DNA-BINDING PROTEIN 1 (CHD-1).//7.5e-41:164:48//HO-MO SAPIENS (HUMAN),//O14646
- F-NT2RP3001727//HYPOTHETICAL 37.7 KD PROTEIN ZK686.3 IN CHROMOSOME III.//1.5e-51:240:41// 10 CAENORHABDITIS ELEGANS //P34669
  - F-NT2RP3001730//SEPTIN 2 HOMOLOG (FRAGMENT).//2.4e-122:267:86//HOMO SAPIENS (HUMAN).// Q14141
  - F-NT2RP3001739//INTESTINAL SODIUM/DICARBOXYLATE COTRANSPORTER (NA(+)/DICARBOXYLATE COTRANSPORTER), //0.99:63:34//RATTUS NORVEGICUS (RAT), //P70545
  - F-NT2RP3001752//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//4.0e-21:60:85//HOMO SAPIENS (HUMAN).//
    - F-NT2RP3001753//HYPOTHETICAL PROTEIN KIAA0127.//7.9e-12:83:44//HOMO SAPIENS (HUMAN).// Q14140
- 20 F-NT2RP3001764//DUAL SPECIFICITY PROTEIN PHOSPHATASE 6 (EC 3.1.3.48) (EC 3.1.3.16) (DUAL SPE-CIFICITY PROTEIN PHOSPHATASE PYST1).//7.7e-25:146:36//HOMO SAPIENS (HUMAN).//Q16828
  - F-NT2RP3001777//SERINE/THREONINE-PROTEIN KINASE STE20 HOMOLOG (EC 2.7.1.-).//0.0096;204;25// CANDIDA ALBICANS (YEAST), //Q92212
- F-NT2RP3001782//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L),//0.91:34:44//PONGO PYGMAEUS ABELII 25 (SUMATRAN ORANGUTAN),//P92694
- F-NT2RP3001792//HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN M (HNRNP M), // 1.8e-33:159:53// HOMO SAPIENS (HUMAN) //P52272
- F-NT2RP3001799//LIGHT-HARVESTING PROTEIN B800/830/1020, ALPHA-2 CHAIN (EHS-ALPHA-2) (ANTEN-NA PIGMENT PROTEIN, ALPHA-2 CHAIN),//0.14:46:28//ECTOTHIORHODOSPIRA HALOCHLORIS.//P80103 F-NT2RP3001819//PROCOLLAGEN ALPHA 2(I) CHAIN PRECURSOR://0.00030:77:36//HOMO SAPIENS (HU-
- 30 MAN).//P08123 F-NT2RP3001844//OCTAMER-BINDING TRANSCRIPTION FACTOR 1 (OTF-1) (NF-A1) (FRAGMENT).//0.99:
  - 43:34//MACROPUS EUGENII (TAMMAR WALLABY), //Q28466 F-NT2RP3001854//FIBRINOGEN- AND IG-BINDING PROTEIN PRECURSOR (MRP PROTEIN).//9.3e-10:213:
  - 24/ISTREPTOCOCCUS PYOGENES //P30141 F-NT2RP3001855//HOMEOBOX PROTEIN PKNOX1 (HOMEOBOX PROTEIN PREP-1).//2.6e-61;220;60//HO-
    - MO SAPIENS (HUMAN).//P55347 F-NT2RP3001857//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//1.0e-13:213:24//PODOSPORA AN-SERINA //O00808
- 40 F-NT2RP3001896//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//0.074:124:34//HOMO SA-PIENS (HUMAN).//Q15428
  - F-NT2RP3001898//REGULATORY PROTEIN E2.//0.36:131:29//CANINE ORAL PAPILLOMAVIRUS (COPV).// O89420
- F-NT2RP3001915//CHITIN BIOSYNTHESIS PROTEIN CHS5 (CAL3 PROTEIN).//0.0021;237;23//SACCHARO-45 MYCES CEREVISIAE (BAKER'S YEAST), //Q12114
- F-NT2RP3001926//HYPOTHETICAL 14.0 KD PROTEIN IN RPL15B-GCR3 INTERGENIC REGION.//1.0:63:34// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q03880
  - F-NT2RP3001929//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!/1.4e-14:35:60//HOMO SAPIENS (HUMAN).// P39195
  - F-NT2RP3001931//HYPOTHETICAL 59.3 KD PROTEIN IN TAP42-ARP9 INTERGENIC REGION.//0.86:162:24// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q05040
    - F-NT2RP3001938//GLYCOPROTEIN GP50.//0.0036:54:40//PSEUDORABIES VIRUS (STRAIN RICE) (PRV).// P07645
- F-NT2RP3001943//33.2 KD PROTEIN IN DIND-RPH INTERGENIC REGION (ORF X).//1.0:113:27//ES-CHERICHIA COLI.//P23839
- F-NT2RP3001944//HYPOTHETICAL 47.6 KD PROTEIN C16C10.5 IN CHROMOSOME III.//4.1e-56:208:47// CAENORHABDITIS ELEGANS.//Q09251
  - F-NT2RP3001969//PUFF II/9-2 PROTEIN PRECURSOR.//0.0078:149:26//SCIARA COPROPHILA (FUNGUS

GNAT).//P22312

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- F-NT2RP3001989//SPERM PROTAMINE P1 (CYSTEINE-RICH PROTAMINE).//1.0:41:31//MUS MUSCULUS (MOUSE).//P02319
- F-NT2RP3002002//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.2e-44:69:79//HOMO SAPIENS (HUMAN).//
- F-NT2RP3002004//TRANSCRIPTION FACTOR BF-2 (BRAIN FACTOR 2) (BF2).//0.00024:45:40//MUS MUSCU-LUS (MOUSE).//061345
- F-NT2RP3002007/TENASCIN PRECURSOR (TN) (HEXABRACHION) (CYTOTACTIN) (NEURONECTIN) (GMEM) (JI) (MIOTENDINOUS ANTIGEN) (GLIOMA-ASSOCIATED-EXTRACELLULAR MATRIX ANTIGEN) (GP 150-225) (TENASCIN-C), //0.21:115:28/IHOMO SAPIENS (HUMAN), //P24821
  - F-NT2RP3002014//HYPOTHETICAL 32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III.//1.7e-25:139:48// CAENORHABDITIS ELEGANS.//Q09232
  - F-NT2RP3002033//ACTIVATOR OF APOPTOSIS HARAKIRI (NEURONAL DEATH PROTEIN DP5).//0.14:65:41// HOMO SAPIENS (HUMAN).//000198
- 75 F-NT2RP3002045/ALPHA-ADAPTIN C (CLATHRIN ASSEMBLY PROTEIN COMPLEX 2 ALPHA-C LARGE CHAIN) (100 KD COATED VESICLE PROTEIN C) (PLASMA MEMBRANE ADAPTOR HA2/AP2 ADAPTIN ALPHA
  - C SUBUNIT).//8.1e-108:192:98/MUS MUSCULUS (MOUSE).//P17427 F-NT2RP3002054//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.046:176:31//STREPTO-MYCES FRADIAE.//P20186
- 20 F-NT2RP3002056//140 KD NUCLEOLAR PHOSPHOPROTEIN (NOPP140).//1.4e-07:245:25//RATTUS NOR-VEGICUS (RAT).//P41777
- F-NT2RP3002057//SMALL HYDROPHOBIC PROTEIN.//1.0:12:66//SIMIAN VIRUS 5 (STRAIN W3) (SV5).//
- F-NT2RP3002062//PROTEASE A INHIBITOR 3 (PROTEINASE INHIBITOR I(A)3).//1.0:49:32//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P01094
  - F-NT2RP3002063//ACYL CARRIER PROTEIN (ACP).//0.99:38:31//HAEMOPHILUS INFLUENZAE.//P43709
    F-NT2RP3002081//HYPOTHETICAL 100.5 KD PROTEIN C1B9.04 IN CHROMOSOME I.//5.8e-35:253:37//
- SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//010429
  F-NT2RP3002097//HYPOTHETICAL 98.1 KD PROTEIN IN SPX19-GCR2 INTERGENIC REGION.//6.2e-06:99:
  31//SACCHAROMYCES CEREVISIAE (BAKERS YEAST)//P40164
- F-NT2RP3002102//HYPOTHETICAL 7.4 KD PROTEIN.//0.68:34:47//THERMOPROTEUS TENAX VIRUS 1
  (STRAIN KRA1) (TTV1).//P19302
  - F-NT2RP3002108//HYPOTHETICAL 105.5 KD PROTEIN R13F6.10 IN CHROMOSOME III.//7.9e-19:179:34// CAENORHABDITIS ELEGANS.//021986
- 35 F-NT2RP3002142//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.6e-17:37:75//HOMO SAPIENS (HUMAN).// P39188
- F-NT2RP3002146//CUTICLE COLLAGEN 40.//0.00034:90:37//CAENORHABDITIS ELEGANS.//P34804
  - F-NT2RP3002147//SALIVARY PROLINE-RICH PROTEIN PO PRECURSOR (ALLELE S).//0.011:166:28//HOMO SAPIENS (HUMAN).//P10163
- 40 F-NT2RP3002151//G1 TO S PHASE TRANSITION PROTEIN 1 HOMOLOG (GTP-BINDING PROTEIN GST1-HS).//4.8e-11:60:53//HOMO SAPIENS (HUMAN).//P15170
  - F-NT2RP3002163/TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII-135) (TAFII-130) (TARII130).//0.028:191:29//HOMO SAPIENS (HUMAN).//O00268
- F-NT2RP3002165//TRANSCRIPTIONAL REGULATOR PROTEIN HCNGP.//2.3e-131:223:91//MUS MUSCULUS (MOUSE).//Q02614
  - F-NT2RP3002168/ID-ALANYL CARRIER PROTEIN (DCP)//1.0:66:33/ILACTOBACILLUS CASEI.//P56163 F-NT2RP3002173//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.4e-26:114:62//HOMO SAPIENS (HU-MAN).///939194
  - F-NT2RP3002181//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.25:31:38//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01645
- F-NT2RP3002244//SPERM PROTAMINE P1 (CYSTEINE-RICH PROTAMINE).//0.069:16:62//OVIS ARIES (SHEEP), AND CAPRA HIRCUS (GOAT).//P04102
  - F-NT2RP3002248//MICROFIBRILLAR-ASSOCIATED PROTEIN 1 (ASSOCIATED MICROFIBRIL PROTEIN) (AMF).//0.0079:187:24//GALLUS GALLUS (CHICKEN).//P55080
- 55 F-NT2RP3002255//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//4.6e-10:168:34//MUS MUSCULUS (MOUSE).//P05143
  - F-NT2RP3002273//SCD6 PROTEIN.//1.5e-11:160:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P45978

F-NT2RP3002276//PROBABLE E4 PROTEIN.//0.91:54:29//HUMAN PAPILLOMAVIRUS TYPE 16.//P06922 F-NT2RP3002303//HYPOTHETICAL. 30.2 KD PROTEIN C4D7.04C IN CHROMOSOME L//1.7e-42:191:43// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//014171 F-NT2RP3002304

F-NT2RP3002330//NNP-1 PROTEIN.//0.52:140:18//MUS MUSCULUS (MOUSE).//P56183

F-NT2RP3002343//5E5 ANTIGEN.//0.0056:189:30//RATTUS NORVEGICUS (RAT).//Q63003

F-NT2RP3002351/NAD-DEPENDENT METHYLENETETRAHYDROFOLATE DEHYDROGENASE (EC 1.5.1.15) / METHENYLIETRAHYDROFOLATE CYCLOHYDROLASE (EC 3.5.4.9) MITOCHONDRIAL PRECURSOR.// 1.0e-66:196.68/HOMO SAPIENS (HUMAN)./P13995

10 F-NT2RP3002352//PRESYNAPTIC PROTEIN SAP102 (SYNAPSE-ASSOCIATED PROTEIN 102) (NEUROEN-DOCRINE-DLG) (NE-DLG).//0,79:173:27//HOMO SAPIENS (HUMAN).//Q92796

F-NT2RP3002377//PUTATIVE HELICASE YGR271W.//1.0e-56:216:44//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53327

F-NT2RP3002399//MINICHROMOSOME MAINTENANCE PROTEIN 6.//1.4e-19:136:31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53091

F-NT2RP3002402//EBNA-6 NUCLEAR PROTEIN (EBNA-3C) (EBNA-4B).//0.74:107:36//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03204

F-NT2RP3002455//DNAJ PROTEIN (FRAGMENT).//5.6e-06:57:42//AGROBACTERIUM TUMEFACIENS.//

F-NT2RP3002484//HYPOTHETICAL 46.5 KD PROTEIN C12B10.04 IN CHROMOSOME 1.//0.00032:52:48// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10438

F-NT2RP3002501//HYPOTHETICAL 34.9 KD PROTEIN IN FRE2-JEN1 INTERGENIC REGION.//9.4e-42:209: 42//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36007

F-NT2RP3002512//HYPOTHETICAL 37.4 KD PROTEIN IN GPM1-MCR1 INTERGENIC REGION.//7.7e-32:162: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36059

F-NT2RP3002529/IPUTATIVE VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN C2G11.03C.//2.1e-45: 241:43/ISCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09805
F-NT2RP3002545

F-NT2RP3002549//HYPOTHETICAL 26.6 KD PROTEIN T19C3.4 IN CHROMOSOME III.//2.8e-41:161:52// CAENORHABDITIS ELEGANS.//Q10010

F-NT2RP3002566//IMMEDIATE-EARLY PROTEIN IE180.//0.56:130:24//PSEUDORABIES VIRUS (STRAIN KA-PLAN) (PRV).//P33479

F-NT2RP3002587

F-NT2RP3002590

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35 F-NT2RP3002602/JPROTEIN DISULFIDE ISOMERASE PRECURSOR (PDI) (EC 5.3.4.1) (THIOREDOXIN- RE-LATED GLYCOPROTEIN 1).//0.00091:111:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P17967 F-NT2RP3002603/JHYPOTHETICAL 14.2 KD PROTEIN IN BLAB 3'REGION.//1.0:65.40//STREPTOMYCES CA-CAOI.//P33864

F-NT2RP3002628//DNAJ-LIKE PROTEIN SLR0093.//2.4e-17:101:44//SYNECHOCYSTIS SP. (STRAIN PCC 68031.//P50027

F-NT2RP3002631//METALLOTHIONEIN-IB (MT-1B).//0.092:36:33://HOMO SAPIENS (HUMAN).//P07438 F-NT2RP3002650//DUALIN.//3.0e-21:184:37//GALLUS GALLUS (CHICKEN).//Q90830

F-NT2RP3002659//PROCOLLAGEN ALPHA 2(I) CHAIN PRECURSOR.//0.00016:223:33//HOMO SAPIENS (HU-

45 F-NT2RP3002660//40S RIBOSOMAL PROTEIN S27A.//0.16:72:31//CAENORHABDITIS ELEGANS.//P37165 F-NT2RP3002663//OXYSTEROL-BINDING PROTEIN.//5.4e-23:168:41//HOMO SAPIENS (HUMAN).//P22059 F-NT2RP3002671//HYPOTHETICAL 124.5 KD PROTEIN IN SKO1-RPL44A INTERGENIC REGION.//6.0e-38: 203:43//SACCHAROMYCES CEREVISIAE (BAKERS YEAST).//P53893

F-NT2RP3002682//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3).//0.25:63:31//ARTEMIA SALINA (BRINE SHRIMP).//P19049

F-NT2RP3002687//HYPOTHETICAL 30.4 KD PROTEIN IN LEF3-IAP2 INTERGENIC REGION.//0.029:60:36// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41469

F-NT2RP3002688//KINESIN-LIKE PROTEIN KIF1B.//5.3e-61:130:88//MUS MUSCULUS (MOUSE).//Q60575 F-NT2RP3002701//SPERM\_MITOCHONDRIAL\_CAPSULE\_SELENOPROTEIN\_(MCS).//7.4e-05:109:33//MUS

MUSCULUS (MOUSE)//P15265
F-NT2RP3002713//PROBABLE ATP-DEPENDENT RNA HELICASE DDX10 (DEAH BOX PROTEIN 10).//0.77:
70:32//HOMO SAPIENS (HUMAN).//0.13206

F-NT2RP3002763//HYPOTHETICAL 11.3 KD PROTEIN C2C6.07 IN CHROMOSOME I.//6.7e-11:66:40//

- SCHIZOSACCHAROMYCES POMBE (FISSION YEAST) //O14056
- F-NT2RP3002770//COLLAGEN ALPHA 1(IX) CHAIN (FRAGMENT).//0.33:87:34//MUS MUSCULUS (MOUSE).// Q05722
- F-NT2RP3002785//LETHAL(2)DENTICLELESS PROTEIN (DTL83 PROTEIN).//9.7e-36:187:39//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q24371
  - F-NT2RP3002799//!!!! ALU SUBFAMILY J WARNING ENTRY!!!!/5.6e-08:41:73//HOMO SAPIENS (HUMAN).//
  - F-NT2RP3002810//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.0034:35:65//HOMO SAPIENS (HUMAN).// P39193
- 10 F-NT2RP3002818//MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENP-B).//3.2e-17:148:37//MUS MUSCULUS (MOUSE).//P27790
  - F-NT2RP3002861//HYPOTHETICAL 70.2 KD PROTEIN IN GSH1-CHS6 INTERGENIC REGION.//1.7e-05:95: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P42951
- F-NTZRP3002869/TRYPSIN INHIBITOR II (BDTI-II).//0.97.23:39/BRYONIA DIOICA (RED BRYONY).//P11968
  15 F-NTZRP3002876/IACIDC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP33).//0.00017:140:31//RATTUS NORVEGICUS (RAT).//P04474
  - F-NT2RP3002877//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.5e-06:55:60//HOMO SAPIENS (HUMAN).//
  - F-NT2RP3002909//P53-BINDING PROTEIN 53BP2 (BCL2-BINDING PROTEIN) (BBP).//4.6e-08:129:38//HOMO SAPIENS (HLIMAN) //013625
  - F-NT2RP3002911//HYPOTHETICAL PROTEIN C18.//0.99:26:50//SWINEPOX VIRUS (STRAIN KASZA) (SPV).// P32217
  - F-NT2RP3002948//RING CANAL PROTEIN (KELCH PROTEIN).//1.2e-23:113:47//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- 25 F-NT2RP3002953//CADHERIN-RELATED TUMOR SUPPRESSOR PRECURSOR (FAT PROTEIN).//0.55:116: 27//DROSOPHILA MELANOGASTER (FRUIT FLY).//P33450
  - F-NT2RP3002955/HYPOTHETICAL 16.5 KD PROTEIN IN BLTR-SPOIIIC INTERGENIC REGION.//0.87:67:37// BACILLUS SUBTILIS.//P54445
- F-NT2RP3002969//LONG-CHAIN-FATTY-ACID-COA LIGASE 4 (EC 6.2.1.3) (LONG-CHAIN ACYL-COA SYN-THETASE 4) (LACS 4).//6.7e-56:189:59//HOMO SAPIENS (HUMAN).//060488
- F-NT2RP3002972//HYPOTHETICAL 73.0 KD PROTEIN IN CLA4-MID1 INTERGENIC REGION.//0.0028:147:27// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P48566
  - F-NT2RP3002978//PROBABLE E5 PROTEIN.//0.15:55:36//HUMAN PAPILLOMAVIRUS TYPE 51.//P26553 F-NT2RP3002985//METALLOTHIONEIN (MT).//0.0031:49:42//PLEURONECTES PLATESSA (PLAICE).//
  - P07216
    F-NT2RP3002988//NEUROGENIC LOCUS NOTCH HOMOLOG PROTEIN 1 PRECURSOR (MOTCH PROTEIN)/1.0·111:28//MUS MUSCULUS (MOUSE)//Q01705
    - F-NT2RP3003008/HYPOTHETICAL 54.7 KD PROTEIN F37A4.1 IN CHROMOSOME III.//0.96:112:25//
- 40 F-NT2RP3003032

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- F-NT2RP3003059//HYPOTHETICAL 52.3 KD PROTEIN C56F8.06C IN CHROMOSOME | PRECURSOR.//9.7e-27:216:37//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10254
- F-NT2RP3003061//ANKYRIN R (ANKYRINS 2.1 AND 2.2) (ERYTHROCYTE ANKYRIN).//3.7e-25:167:34//HOMO SAPIENS (HUMAN).//P16157
- 45 F-NT2RP3003068//SERYL-TRNA SYNTHETASE (EC 6.1.1.11) (SERINE-TRNA LIGASE) (SERRS) (FRAG-MENT).//0.074:82:39//SULFOLOBUS SOLFATARICUS.//O33780
  - F-NT2RP3003071/VASODILATOR-STIMULATED PHOSPHOPROTEIN (VASP).//0.0085:128:30//HOMO SAPIENS (HUMAN).//P50552
  - F-NT2RP3003078//SPERM ACROSOMAL PROTEIN FSA-ACR.1 PRECURSOR (FRAGMENT).//0.028:165:31// VULPES VULPES (RED FOX).//P53353
- F-NT2RP3003101//TETRACYCLINE RESISTANCE PROTEIN, CLASS C (TETA(C)).//1.0e-14:243:25//ES-CHERICHIA COLI.//P02981
  - F-NT2RP3003121//SUPPRESSOR PROTEIN SRP40://7.4e-05:143:27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)://P32583
- F-NT2RP3003133//65 KD YES-ASSOCIATED PROTEIN (YAP65),//0.024:61:42//GALLUS GALLUS (CHICK-EN),//P46936
  - F-NT2RP3003138//KINESIN-LIKE PROTEIN KIF4.//1.1e-118:151:93//MUS MUSCULUS (MOUSE).//P33174
    F-NT2RP3003139//ATP-BINDING CASSETTE TRANSPORTER ABC1.//1.0:70:30//SCHIZOSACCHAROMYCES

POMBE (FISSION YEAST) //Q92337

F-NT2RP3003145//MILK FAT GLOBULE-EGF FACTOR 8 PRECURSOR (MFG-E8) (HMFG) (BREAST EPITHE-LIAL ANTIGEN BA46) (MFGM).//2.0e-12:121:37//HOMO SAPIENS (HUMAN).//Q08431 F-NT2RP3003150

F-NT2RP3003157//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//4.0e-79:260:54//HOMO SA-PIENS (HUMAN).//P51522

F-NT2RP3003185//TROPOMYOSIN.//0.077:122:27//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//

F-NT2RP3003193//ZINC FINGER PROTEIN 135.//7.2e-91:239:65//HOMO SAPIENS (HUMAN).//P52742

F-NT2RP3003197//HYPOTHETICAL 28.1 KD PROTEIN IN SIPU-PBPC INTERGENIC REGION.//1.3e-07:117: 34//BACILLUS SUBTILIS.//P42966

F-NT2RP3003203//HYPOTHETICAL 33.5 KD PROTEIN C1D4.02C IN CHROMOSOME I.//9.9e-23:132:39// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10149

F-NT2RP3003204//RAS-LIKE PROTEIN RASB.//0.92:103:27//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P32252

F-NT2RP3003210/VERY HYPOTHETICAL 13.2 KD PROTEIN IN PTC3-SAS3 INTERGENIC REGION.//0.23: 106:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38190

F-NT2RP3003212//SUPPRESSOR PROTEIN SRP40.//0.019:171:23//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583

F-NTZRP3003230//CORONN-LIKE PROTEIN P57.//8.3e-74-183:73//BOS TAURUS (BOVINE).//IOS2176 F-NTZRP3003242//STANNIOCALCIN PRECURSOR.//1.4e-21:127:37//HOMO SAPIENS (HUMAN).//P52823 F-NTZRP300325/I/DOWN REGULATORY PROTEIN OF INTERLEUKIN Z RECEPTOR.//3.1e-51:198:52//MUS MUSCULUS (MOUSE).//P15533 F-NTZRP3003264//E9 PROTEIN/10:31:41//HUMAN PAPILLOMAVIRUS TYPE 48.//080920

25 F-NT2RP3003278//45.8 KD PROTEIN IN SHM1-MRPL37 INTERGENIC REGION.//8.6e-07:80:33//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P38344

F-NT2RP3003282//DYNAMIN 2 (DYNAMINI UDMN)//8.0e-108:226:88//MUS MUSCULUS (MOUSE).//P39054 F-NT2RP3003290//BIOH PROTEIN.//0.0055:107:30//ESCHERICHIA COLL//P13001 F-NT2RP3003301//MITOCHONDRIAL LON PROTEASE HOMOLOG 1 PRECURSOR (EC 3.4.21.-).//1.3e-69:

200:55//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS)//084948 F-NTZRP3003302//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG//6.4e-69:102:66//HOMO SAPIENS (HU-MAN) //P08547

F-NT2RP3003311//MYOSIN II HEAVY CHAIN, NON MUSCLE.//0.18:225:26//ACANTHAMOEBA CASTELLANII (AMOEBA //P)05659

35 F-NT2RP3003313//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.0014:142:33//HOMO SAPIENS (HUMAN).//P10162

F-NT2RP3003327//52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS-A)) (RO(SS-A)) (RO52).//8.8e-18:94:43//MUS MUSCULUS (MOUSE).//Q62191

F-NT2RP3003330//HYPOTHETICAL PROTEIN KIAA0176 (FRAGMENT).//1.3e-20:123:44//HOMO SAPIENS (HUMAN).//014681

F-NT2RP3003344//HYPOTHETICAL 8.8 KD PROTEIN IN ICDC-MINE INTERGENIC REGION.//1.0:28:42//ES-CHERICHIA COLI.//P75991

F-NT2RP3003346//!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//6.9e-26:74:78//HOMO SAPIENS (HU-MAN) //P30191

45 F-NT2RP3003353//HYPOTHETICAL 52.4 KD PROTEIN R08D7.2 IN CHROMOSOME III.//3.7e-10:118:33// CAENORHABDITIS ELEGANS.//P30641

F-NT2RP3003377//PUTATIVE CUTICLE COLLAGEN F09G8.6.//1.5e-05:102:37//CAENORHABDITIS ELE-GANS.//P34391

F-NT2RP3003384 F-NT2RP3003385//SKD3 PROTEIN.//5.1e-83:210:69//MUS MUSCULUS (MOUSE).//Q60649

F-NT2RP3003409//SOX-22 PROTEIN.//0.042:173:28//HOMO SAPIENS (HUMAN).//O15370

F-NT2RP3003411//PROBABLE E3 PROTEIN.//0.17:91:31//BOVINE PAPILLOMAVIRUS TYPE 2.//P11300 F-NT2RP3003427//HOLOTRICIN 3 PRECURSOR.//0.012:36:41//HOLOTRICHIA DIOMPHALIA.//Q25055

55 F-NT2RP3003433 F-NT2RP3003464//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.0042:110: 40/SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214

F-NT2RP3003490

F-NT2RP3003403

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- F-NT2RP3003491//10 KD CHAPERONIN (PROTEIN CPN10) (PROTEIN GROES) (HEAT SHOCK 10 KD PROTEIN) //0 99/49/34// EPTOSPIRA INTERROGANS //P35472
- F-NT2RP3003500//SCY1 PROTEIN.//6.8e-14:192:26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
- F-NT2RP3003543//COLLAGEN ALPHA 5(IV) CHAIN PRECURSOR.//0.0026:175:30//HOMO SAPIENS (HU-MAN).//P29400
  - F-NT2RP3003552//ANNEXIN VII (SYNEXIN) (FRAGMENT).//0.19:21:47//BOS TAURUS (BOVINE).//P20072 F-NT2RP3003555//HYPOTHETICAL 3.2.6 KD PROTEIN IN MET30-PIG2 INTERGENIC REGION.//T.3e-27:159: 43//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40529
- F-NT2RP3003564//RNA REPLICASE POLYPROTEIN (EC 2.7.7.48).//1.0:99:30//TURNIP YELLOW MOSAIC VI-RUS //P10358
  - F-NT2RP3003572//PUTATIVE CUTICLE COLLAGEN F09G8.6.//0.33:128:32//CAENORHABDITIS ELEGANS.// P34391
- F-NT2RP3003576//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//7.1e-28:58:77//HOMO SAPIENS (HUMAN).//
  15 P39195
  - F-NTZRP3003588//RAS-RELATED PROTEIN RAB-10.//5.4e-54:114:94//CANIS FAMILIARIS (DOG).//P24409
    F-NTZRP3003588//ROAGULATION FACTOR XII PRECURSOR (EC 3.4.21.38) (HAGEMAN FACTOR) (HAF).//
    2.0e-15:88-40//HOMO SAPIENS (HUMAN).//P00748
- F-NT2RP3003625/MALE SPECIFIC SPERM PROTEIN MST84DD.//0.99:22:50//DROSOPHILA MELA-20 NOGASTER (FRUIT FLY).//Q01645
- F-NT2RP3003656//HOMEOBOX PROTEIN OTX3 (ZOTX3).//0.30:111:25//BRACHYDANIO RERIO (ZE-BRAFISH) (ZEBRA DANIO).//090267
  - F-NT2RP3003659//HYPOTHETICAL 49.8 KD PROTEIN IN RPL14B-GPA1 INTERGENIC REGION.//1.1e-20:127: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38755
- 25 F-NT2RP3003665//PENAEIDIN-3C PRECURSOR (P3-C).//0.34:52:34//PENAEUS VANNAMEI (PENOEID SHRIMP) (JEROPEAN WHITE SHRIMP).//P61060 F-NT2RP3003672//T-CELL SURFACE GLYCOPROTEIN E2 PRECURSOR (E2 ANTIGEN) (CD99) (MIC2 PRO-
  - TEIN) (12E7)//8.76-15:146:42//HOMO SAPIENS (HUMAN).//P14209
    F-NT2R79003860//HYPOTHETICAL, 55.1 KD PROTEIN IN FAB1-PES4 INTERGENIC REGION.//4.3e-25:159:
- 40//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43601
  - F-NT2RP3003686//NONHISTONE CHROMOSOMAL PROTEIN HMG-17.//0.067:63:31//GALLUS GALLUS (CHICKEN).//P02314
    - F-NT2RP3003701//F-SPONDIN PRECURSOR.//1.8e-13:193:27//RATTUS NORVEGICUS (RAT).//P35446 F-NT2RP3003716//SLIT PROTEIN PRECURSOR.//1.3e-12:150:34//DROSOPHILA MELANOGASTER (FRUIT
- 35 FLY).//P24014 F-NT2RP3003726//INSERTION ELEMENT IS136 HYPOTHETICAL 16.9 KD PROTEIN).//0.47:109:28//AGRO-
  - BACTERIUM TUMEFACIENS.//P05680 F-NT2RP3003746//HYPOTHETICAL 7.7 KD PROTEIN IN FIXX 3'REGION (ORF1).//0.57:34:38//AZORHIZO-
  - BIUM CAULINODANS.//P26486 F-NT2RP3003795///III ALU SUBFAMILY SX WARNING ENTRY !!!//4 3e-10:40:90//HOMO SAPIENS (HUMAN) //
- F-NT2RP3003795/!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//4.3e-10:40:90/iHOMO SAPIENS (HUMAN).// P39195 F-NT2RP3003799/MATING-TYPE PHEROMONE BBP1(3) PRECURSOR./i0.75:60:36//SCHIZOPHYLLUM
  - COMMUNE (BRACKET FUNGUS).//P78744

    F-NT2RP3003800//PROTO-ONCOGENE TYROSINE-PROTEIN KINASE SRC (EC 2.7.1.112) (P60-SRC).//4.2e-
- 45 51:72:95//GALLUS (CHICKEN).//P00523 F-NT2RP3003005//HYPOTHETICAL 32.1 KD PROTEIN IN DBP7-GCN3 INTERGENIC REGION.//0.00069:160:
  - 25//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36121
    F-NT2RP3003809//COLLAGEN ALPHA 1(III) CHAIN (FRAGMENTS).//0.028:135:35//GALLUS GALLUS (CHICK-
  - EN),//P12105
    F-NT2RP3003819//C-HORDEIN (PCP387) (FRAGMENT),//0.0026:90:33//HORDEUM VULGARE (BARLEY),//
- P06472

- F-NT2RP3003825//PHOSPHATIDYLCHOLINE TRANSFER PROTEIN (PC-TP).//5.6e-20:174:31//BOS TAURUS (BOVINE).//P02720
- F-NT2RP3003828//ADENYLATE CYCLASE, TYPE V (EC 4.6.1.1) (ATP PYROPHOSPHATE-LYASE) (CA(2+)55 INHIBITABLE ADENYLYL CYCLASE).//0.0017:111:38//CANIS FAMILIARIS (DOG).//P30803
  - F-NT2RP3003831//ENDONUCLEASE G PRECURSOR (EC 3.1.30.-) (ENDO G).//1.1e-37:187:42//MUS MUSCU-LUS (MOUSE).//008600
    - F-NT2RP3003833//HYPOTHETICAL 6.4 KD PROTEIN IN INTE-PIN INTERGENIC REGION.//1.0:38:39//ES-

- CHERICHIA COLL//P75979
- F-NT2RP3003842

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- F-NT2RP3003846//RETINAL DEGENERATION B PROTEIN (PROBABLE CALCIUM TRANSPORTER RDGB).// 0.61:54:35//DROSOPHILA MELANOGASTER (FRUIT FLY).//P43125
- F-NT2RP3003870//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.83:51:37//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643
  - F-NT2RP3003876//PROTEIN TRANSPORT PROTEIN SEC2.//0.0017:151:27//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P17065
  - F-NT2RP3003914//UDP-GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (EC 2.4.1.-) (DUGT).//3.3e-23:76:64//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q09332
  - F-NT2RP3003918//VESICLE-ASSOCIATED MEMBRANE PROTEIN/SYNAPTOBREVIN BINDING PROTEIN (VAP-33),//5.5e-45:127:69//APLYSIA CALIFORNICA (CALIFORNIA SEA HARE),//Q16943 F-NT2RP3003932
  - F-NT2RP3003989//PREPROTEIN TRANSLOCASE SECE SUBUNIT //0 96:46:32//THERMOTOGA MARITIMA // P35874
  - F-NT2RP3003992//NUCLEAR LOCALIZATION SEQUENCE BINDING PROTEIN (P67).//0.0011:170:26//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P27476
    - F-NT2RP3004013//DOUBLE-STRANDED RNA-SPECIFIC EDITASE 1 (EC 3.5.-.) (DSRNA ADENOSINE DEAM-INASE) (RNA EDITING ENZYME 1).//3.6e-21:134:45//RATTUS NORVEGICUS (RAT).//P51400
- 20 F-NT2RP3004016//HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION.//0.00021:64:40// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41479
  - F-NT2RP3004041//SPERM PROTAMINE P1.//0.0028:43:46//ORNITHORHYNCHUS ANATINUS (DUCKBILL PLATYPUS) //P35307
- F-NT2RP3004051//MICROBIAL COLLAGENASE PRECURSOR (EC 3.4,24.3) (120 KD COLLAGENASE),// 25 0.0079:194:24//CLOSTRIDIUM PERFRINGENS.//P43153
  - F-NT2RP3004070//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.4e-11:51:72//HOMO SAPIENS (HUMAN).// P39188
  - F-NT2RP3004078//DNA BINDING PROTEIN RFX2.//2.7e-114:243:87//MUS MUSCULUS (MOUSE).//P48379 F-NT2RP3004093//HYPOTHETICAL 32.3 KD PROTEIN IN RHSE-NARV INTERGENIC REGION (ORFB).//8.0e-13:111:41//ESCHERICHIA COLI.//P37757
- F-NT2RP3004095//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/1.5e-17:72:65//HOMO SAPIENS (HUMAN).// P39188
  - F-NT2RP3004110//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.6e-10:51:72//HOMO SAPIENS (HUMAN).// P39195
- 35 F-NT2RP3004125//ZINC FINGER PROTEIN 75.//1.1e-28:118:47//HOMO SAPIENS (HUMAN).//P51815 F-NT2RP3004145//AEROLYSIN REGULATORY PROTEIN.//0.012;45;33//AEROMONAS SOBRIA.//P09165 F-NT2RP3004148//METALLOTHIONEIN-I (MT-1).//0.055:18:50//COLUMBA LIVIA (DOMESTIC PIGEON).//
  - F-NT2RP3004155//UBIQUINONE BIOSYNTHESIS PROTEIN COQ7 HOMOLOG.//1.7e-82:178:89//RATTUS
- 40 NORVEGICUS (RAT).//Q63619 F-NT2RP3004189//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//1.7e-11:215:24//PODOSPORA AN-
  - F-NT2RP3004206//CROOKED NECK PROTEIN.//3.8e-101:241:73//DROSOPHILA MELANOGASTER (FRUIT FLY) //P17886
- 45 F-NT2RP3004207//CUTICLE COLLAGEN 12 PRECURSOR.//0.13:130:33//CAENORHABDITIS ELEGANS.//
  - F-NT2RP3004209//UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 4 (EC 3.1.2.15) (UBIQUITIN THIOLESTE-RASE 4) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 4) (DEUBIQUITINATING ENZYME 4) (UBIQUI-TOUS NUCLEAR PROTEIN HOMOLOG).//6.5e-16:207:29//HOMO SAPIENS (HUMAN).//Q13107
  - F-NT2RP3004215//PROTEIN TRANSPORT PROTEIN SEC61 GAMMA SUBUNIT.//1.0:69:31//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST),//P35179
    - F-NT2RP3004242//HYPOTHETICAL 30.2 KD PROTEIN ZK632.12 IN CHROMOSOME III.//1.1e-64:191:63// CAENORHARDITIS ELEGANS //P34657
- F-NT2RP3004246//RING3 PROTEIN (KIAA9001).//0.060:101:28//HOMO SAPIENS (HUMAN).//P25440
- 55 F-NT2RP3004253//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//1.1e-07:184:35//BOS TAURUS (BOVINE).// P02453
  - F-NT2RP3004258//SUPPRESSOR PROTEIN SRP40.//4.9e-08:98:39//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32583

- F-NT2RP3004262//DNAJ PROTEIN HOMOLOG 1 (HDJ-1) (HEAT SHOCK PROTEIN 40) (HSP40).//1.6e-63:210: 61//HOMO SAPIENS (HUMAN).//P25685
- F-NT2RP3004282//HYPOTHETICAL PROTEIN F44G4.1 IN CHROMOSOME II (FRAGMENT).//1.6e-29:177:38// CAENORHABDITIS ELEGANS.//P54073
- F-NT2RP3004332//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RPB1) (FRAG-MENT)./0.030-118:36//CRICETULUS GRISEUS (CHINESE HAMSTER).//P11414 F-NT2RP3004334
  - F-NT2RP3004341//ALPHA-INTERNEXIN (ALPHA-INX).//0.91:110:26//MUS MUSCULUS (MOUSE).//P46660
  - F-NT2RP3004348//HYPOTHETICAL 105.3 KD PROTEIN C01G6.5 IN CHROMOSOME III.//0.60:198:24// CAENORHABDITIS ELEGANS.//P46012

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- F-NT2RP3004349//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.0e-37:60:76//HOMO SAPIENS (HUMAN).// P39193
- F-NT2RP3004378//HYPOTHETICAL 18.8 KD PROTEIN IN GNTR-GGT INTERGENIC REGION (O162).//0.0026: 76:28//ESCHERICHIA COLI.//P46854
- 15 F-NT2RP3004399//LEUCINE-RICH PRIMARY RESPONSE PROTEIN 1 (FOLLICLE-STIMULATING HORMONE PRIMARY RESPONSE PROTEIN).//4.4e-109:212:96//HOMO SAPIENS (HUMAN).//Q92674
  - F-NT2RP3004424//1TV-1 PROTEIN.//4.5e-18:60:70//HOMO SAPIENS (HUMAN).//C13155 F-NT2RP3004428// METALLOTHIONEIN-A (MTA).//0.0010:36:47//STRONGYLOCENTROTUS PURPURATUS (PURPLE SEA UR-CHIN).//PQ4734
- F-NT2RP3004451//MYOSIN IC HEAVY CHAIN.//0.00072:113:34//ACANTHAMOEBA CASTELLANII (AMOE-BA).//P10569
  - F-NT2RP3004454//VERPROLIN.//3.3e-07:156:29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
- F-NT2RP3004466//HYPOTHETICAL PROTEIN F-215.//0.0013:125:32//HUMAN ADENOVIRUS TYPE 2.//
  P03291
  - F-NT2RP3004470//HYPOTHETICAL 15.4 KD PROTEIN C16C10.11 IN CHROMOSOME III.//1.0:33:51// CAENORHABDITIS ELEGANS.//Q09254
    - F-NT2RP3004472//GERM CELL-LESS PROTEIN.//7.3e-33:170:40//DROSOPHILA MELANOGASTER (FRUIT FLY) //001820
- 30 F-NT2RP3004475//RHO-GAP HEMATOPOIETIC PROTEIN C1 (P115) (KIAA0131).//8.4e-54:214:46//HOMO SA-PIENS (HUMAN).//P98171
  - F-NT2RP3004480//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS35.//3.9e-47:199:49//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P34110
  - F-NT2RP3004490//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//0.0013:121:33// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437
  - F-NT2RP3004498/HYPOTHETICAL 43.5 KD PROTEIN IN COTD-KDUD INTERGENIC REGION PRECUR-SOR //0.066.87:35/BACILLUS SUBTILIS //P50840 F-NT2RP3004503/IIIII ALU SUBFAMILY SQ WARNING ENTRY !!!!/2.0e-34:102:69/HOMO SAPIENS (HU-
  - MAN).//P39194
    F-NT2RP3004504//SUPPRESSOR\_PROTEIN\_SRP40.//0.64:93:34//SACCHAROMYCES\_CEREVISIAE\_(BAK-
  - ER'S YEAST),//P32583

    F-NT2RP3004507//MOB1 PROTEIN (MPS1 BINDER 1).//Z.2e-16:90:42//SACCHAROMYCES CEREVISIAE
    - F-N12RF30U49011MU51 PROTEIN (MPS1 BINDER 1).112.28-10:30:42//SACCHAROMTOES CEREVISIA (BAKER'S YEAST).1/P40484
      F-N17RF30014527
- 45 F-NT2RP3004534//S-PHASE ENTRY CYCLIN 6.//0.38:148:22//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32943
  - F-NT2RP3004539//INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN 1 PRECURSOR (IGFBP-1) (IBP-1) (IGF-BINDING PROTEIN 1).//0.38:89:38//RATTUS NORVEGICUS (RAT).//P21743
  - F-NT2RP3004544//CYTADHERENCE HIGH MOLECULAR WEIGHT PROTEIN 2 (CYTADHERENCE ACCESSO-RY PROTEIN 2).//0.0024:200:24//MYCOPLASMA PNEUMONIAE.//P75471
- F-NT2RP3004566//GASTRULA ZINC FINGER PROTEIN XLCGF17.1 (FRAGMENT).//4.6e-25:126:43//XENO-PUS LAEVIS (AFRICAN CLAWED FROG).//P18713
  - F-NT2RP3004569//ANKYRIN.//8.3e-07:150:28//MUS MUSCULUS (MOUSE),//Q02357
- F-NT2RP3004572/TRANSCRIPTION INITIATION FACTOR TFIID 150 KD SUBUNIT (TAFII-150) (TAFII150).//
  1.6e-70:247:54//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q24325
- F-NT2RP3004578//CENTROMERIC PROTEIN E (CENP-E PROTEIN).//1.5e-10:210:26//HOMO SAPIENS (HU-MAN).//Q02224
  - F-NT2RP3004594//P54 PROTEIN PRECURSOR.//0.0044:230:24//ENTEROCOCCUS FAECIUM (STREPTO-

COCCUS FAECIUM) //P13692

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- F-NT2RP3004617//DOWN REGULATORY PROTEIN OF INTERLEUKIN 2 RECEPTOR://1.5e-14:113:34//MUS MUSCULUS (MOUSE).//P15533
- F-NT2RP3004618//HYPOTHETICAL 115.4 KD PROTEIN ZK757.3 IN CHROMOSOME III.//4.5e-08:149:30// CAENORHABDITIS ELEGANS.//P34681
- F-NT2RP3004669//ETHANOLAMINE KINASE (EC 2.7.1.82) (EASILY SHOCKED PROTEIN).//1.0e-24:75:48// DROSOPHILA MELANOGASTER (FRUIT FLY).//P54352
- F-NT2RP3004670//CUTICLE COLLAGEN 21/0.00090:159:29//CAENORHABDITIS ELEGANS.//P17656 F-NT2RP4000008//CHLORINE CHANNEL PROTEIN P64.//4.0e-79:243:62//BOS TAURUS (BOVINE).//P35526
- 10 F-NT2RP4000023
  - F-NT2RP4000035//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.6e-06:46:67//HOMO SAPIENS (HUMAN).// P39194
  - F-NT2RP4000049//CALDESMON (CDM).//0.41:63:34//GALLUS GALLUS (CHICKEN).//P12957
  - F-NT2RP4000051//DUALIN.//2.3e-23:195:37//GALLUS GALLUS (CHICKEN).//Q90830
- 15 F-NT2RP4000078/IRING CANAL PROTEIN (KELCH PROTEIN)./I1.2e-24:182:31/DROSOPHILA MELA-NOGASTER (FRUIT FLY)./I/Q04652 F-NT2RP4000102/IXPAR7 PROTEIN./I1.0:54:33/BACILLUS LICHENIFORMIS./I/Q99166
  - F-NT2RP4000109//SLIT PROTEIN PRECURSOR.//1.9e-60:230:46//DROSOPHILA MELANOGASTER (FRUIT
  - F-N12RF4000109/ISELT PROTEIN PRECORSOR.//1.96-60:230:46//DROSOPHILA MELANOGASTER (PROTEIN FLY).//P24014
- F-NT2RP4000111//CLEAVAGE AND POLYADENYLATION SPECIFICITY FACTOR, 100 KD SUBUNIT (CPSF 100 KD SUBUNIT).//1.4e-91:157:100//BOS TAURUS (BOVINE).//Q10568
  - F-NT2RP4000129//5E5 ANTIGEN.//0.00072:124:37//RATTUS NORVEGICUS (RAT).//Q63003
  - F-NT2RP4000147//ZINC FINGER PROTEIN GCS1.//1.5e-26:119:43//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P35197
- 25 F-NT2RP4000150 F-NT2RP4000151/HYPOTHETICAL 31.0 KD PROTEIN R107.2 IN CHROMOSOME III.//4.2e-31:180:47// CAENORHABDITIS ELEGANS //P32740
  - F-NT2RP4000159//SPORE COAT PROTEIN SP96.//0.84:107:28//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD) //P14328
- 30 F-NT2RP4000167//HYPOTHETICAL 98.1 KD PROTEIN IN SPX19-GCR2 INTERGENIC REGION.//2.4e-08:133: 32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40164
  - F-NT2RP4000185/TRANS-ACTING TRANSCRIPTIONAL PROTEIN ICP0 (VMW118 PROTEIN).//5.4e-05:143: 32//HERBES SIMPLEX VIRUS (TYPE 2 / STRAIN HG52).//P28284
  - F-NT2RP4000210//PAIRED AMPHIPATHIC HELIX PROTEIN.//1.8e-40:258:35//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P22579
    - F-NT2RP4000212/ATRIAL GLAND-SPECIFIC ANTIGEN PRECURSOR (AGSA).//1.4e-20:104:40//APLYSIA CALIFORNICA (CALIFORNICA SEA HARE).//P15287 F-NT2RP4000214//FERREDOXIN.//10.1942/I/MOORELLA THERMOACETICA (CLOSTRIDIUM THERMOACE-
- TICUM).//P00203

  F-NT2RP4000218///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.7e-15:48:60//HOMO SAPIENS (HUMAN).//
- F-N12RP4000218///// ALU SUBPAMILY J WARNING ENTRY (#/////./6-15:48:60//HOMO SAPIENS (#///MAN). P39188 F-NT2RP4000243//DUALIN.//5.8e-78:192:70//GALLUS GALLUS (CHICKEN).//Q90830
  - F-NT2RP4000246/NPC DERIVED PROLINE RICH PROTEIN 1 (NDPP-1).//3.1e-83:207:76//MUS MUSCULUS (MOUSE).//Q03173
- 45 F-NT2RP4000259//GLUTATHIONE PEROXIDASE 2 (EC 1.11.1.9).//5.5e-29:153:43//HELIANTHUS ANNUUS (COMMON SUNFLOWER).//O23968
  - F-NT2RP4000283/IANNEXIN VII (SYNEXIN) (FRAGMENT).//0.9842:40//BOS TAURUS (BOVINE).//P20072 F-NT2RP4000290//HYPOTHETICAL 116.5 KD PROTEIN C20G8.09C IN CHROMOSOME I.//3.5e-71:209:66// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//PB7115
  - F-NT2RP4000312//RAS SUPPRESSOR PROTEIN 1 (RSU-1) (RSP-1 PROTEIN) (RSP-1).//8.9e-22:166:37//HO-MO SAPIENS (HUMAN).//Q15404
    - F-NT2RP4000321//VERPROLIN.//0.00018:260:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P37370
- F-NT2RP4000323//ANTHOPLEURIN B (TOXIN AP-B).//0.42:15:46//ANTHOPLEURA XANTHOGRAMMICA (GI-ANT GREEN SEA ANEMONE).//P01531
  - F-NT2RP4000355//HYPOTHÉTICAL 90.9 KD PROTEIN IN GCN20-CMK1 INTERGENIC REGION.//0.75:125:29// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43596
    - F-NT2RP4000360//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT).//0.27:92:

33//RATTUS NORVEGICUS (RAT) //P10164

- F-NT2RP4000367//HYPOTHETICAL 7.3 KD PROTEIN IN 100 KD PROTEIN REGION.//0.99:52:32//HUMAN AD-FNOVIRUS TYPE 41 //P23691
- F-NT2RP4000370//MITOCHONDRIAL PEPTIDE CHAIN RELEASE FACTOR 1 PRECURSOR (MRF-1).//4.1e-40: 163:52//HOMO SAPIENS (HUMAN).//O75570
- F-NT2RP4000376//PHOSPHOLIPASE A-2-ACTIVATING PROTEIN (PLAP).//4.2e-59:125:80//RATTUS NOR-VEGICUS (RAT) //P54319
  - F-NT2RP4000381//NEUROFILAMENT TRIPLET H PROTEIN (200 KD NEUROFILAMENT PROTEIN) (NF-H).//
    0.00058:194:30//MUS MUSCULUS (MOUSE).//P19246
- F-NT2RP4000398//ZINC FINGER PROTEIN 184 (FRAGMENT).//1.2e-45:153:39//HOMO SAPIENS (HUMAN).// O99676
  - F-NT2RP4000415//HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR://
    0.00066:201:271/SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47179
  - F-NT2RP4000417//PROCESSING ALPHA-1,2-MANNOSIDASE (EC 3.2.1.-) (ALPHA-1,2-MANNOSIDASE 1B).//
    1.8e-25:196:40//MUS MUSCULUS (MOUSE).//P39098
- F-NT2RP4000424//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.0e-15:72:61//HOMO SAPIENS (HUMAN).//
  - F-NT2RP4000448//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//7.0e-23:63:82//HOMO SAPIENS (HUMAN).//
- F-NT2RP4000449//REGULATORY PROTEIN SIR2 (SILENT INFORMATION REGULATOR 2).//1.3e-41:102:45// KLUYVEROMYCES LACTIS (YEAST).//P33294
- F-NT2RP4000455//HOMEOBOX PROTEIN SAX-1 (CHOX-3) (FRAGMENT).//0.00014:92:30//GALLUS GALLUS (CHICKEN).//P19601
- F-NTZRP4000457/JUBIQUITIN CARROXYL-TERMINAL HYDROLASE 7 (EC. 3. 1.2.15) (UBIQUITIN THIOLESTE-RASE 7) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 7) (DEUBIQUITINATING ENZYME 7) (HERPESVI-RUS ASSOCIATED UBIQUITIN-SPECIFIC PROTEASE) //1.0e-29:218:38/HOMO SAPIENS (HUMAN).//093009 F-NTZRP4000480/TRANSCRIPTIONAL REGULATORY PROTEIN ALGP (ALGINATE REGULATORY PROTEIN ALGR3/JV0.049:1172-99/PSELIDOMONAS AFRUIGINOSA //P15/276
  - F-NT2RP4000481//HYPOTHETICAL HELICASE C28H8.3 IN CHROMOSOME III.//2.3e-05:152:23//
    CAENORHABDITIS ELEGANS.//Q09475
- F-NT2RP4000498/IMOB1 PROTEIN (MPS1 BINDER 1).//2.3e-48:172:52//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40484
  - F-NT2RP4000500//HYPOTHETICAL 83.6 KD PROTEIN R05D3.2 IN CHROMOSOME III.//1.3e-23:165:35// CAENORHABDITIS ELEGANS.//P34535
- 35 F-NT2RP4000515//PHOSPHODIESTERASE I (EC 3.1.4.1) (5'-EXONUCLEASE) (5'-NUCLEOTIDE PHOS-PHODIESTERASE) (FRAGMENT).//1.0:48:37//BOS TAURUS (BOVINE).//P15396
  - F-NT2RP4000517//METALLOTHIONEIN-LIKE PROTEIN TYPE 2.//1.0:41:36//VICIA FABA (BROAD BEAN).// Q41657
- F-NT2RP4000518//ATP-DEPENDENT RNA HELICASE ROK1.//1.1e-11:93:36//SACCHAROMYCES CEREVI-49 SIAE (BAKER'S YEAST).//P45818
  - F-NT2RP4000519//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.68:55:40//BOS TAURUS (BOVINE).//
  - F-NT2RP4000524//IGA FC RECEPTOR PRECURSOR (BETA ANTIGEN) (B ANTIGEN).//0.37:187:24//STREP-TOCOCCUS AGALACTIAF //P27951
- 45 F-NT2RP4000528//NPL4 PROTEIN.//2.1e-45:305:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P33755
  - F-NT2RP4000541//HOMEOBOX PROTEIN CHOX-1 (FRAGMENT).//0.23:28:50//GALLUS GALLUS (CHICK-EN).//P13544
    - F-NT2RP4000556//HYPOTHETICAL 34.1 KD PROTEIN C40H1.4 IN CHROMOSOME III.//4.3e-14:174:34//
      CAENORHABDITIS FLEGANS //003574
- O CAENORHABDITIS ELEGANS.//003574 F-NT2RP4000560//HYPOTHETICAL 68.7 KD PROTEIN ZK757.1 IN CHROMOSOME III.//2.1e-19:155:36// CAENORHABDITIS ELEGANS.//P24679
  - F-NT2RP4000588//HYPOTHETICAL PROTEIN E-115.//0.014:64:35//HUMAN ADENOVIRUS TYPE 2.//P03290
- F-NT2RP4000814//SPLICING FACTOR, ARGININE/SERINE-RICH 2 (SPLICING FACTOR SCSS) (SC-35)
  (SPLICING COMPONENT, 35 KD) (PR264 PROTEIN) /Z-P27:1884/GALLUS GALUS (CHICKEN)//P30352
  F-NT2RP4000638//EARLY NODULIN 55-1 PRECURSOR (N-55-1) (FRAGMENT)//I0.55-40-40//GLYCINE MAX
  (SOYPEAN)//GD554
  - F-NT2RP4000648//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2,4e-06:31:74//HOMO SAPIENS (HUMAN).//

P39188

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F-NT2RP46000657//HYPOTHETICAL PROTEIN MJ1065.//2.5e-40:237:40//METHANOCOCCUS JANNAS-CHII //OSR465

F-NT2RP4000704

F-NT2RP4000713//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//4.0e-07:134:40//STREP-TOMYCES FRADIAE.//P20186

F-NT2RP4000724//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE].//1.1e-62:109:88//HOMO SAPIENS (HUMAN).//P10266

F-NT2RP4000728//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.0033:190:25//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323

F-NT2RP4000737//PTB-ASSOCIATED SPLICING FACTOR (PSF),//1.0e-05:114:34//HOMO SAPIENS (HU-MAN),//P23246

F-NT2RP4000739//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:20:50//ANAS PLATYRHYNCHOS (DOMESTIC DUCK).//P50655

F-NT2RP4000781//hYPOTHETICAL 27.7 KD PROTEIN IN CPT1-SPC98 INTERGENIC REGION.//0.0013:67: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53915

F-NT2RP4000787//POLLEN SPECIFIC PROTEIN SF3.//1.3e-13:79:39//HELIANTHUS ANNUUS (COMMON SUNFLOWER).//P29675

F-NT2RP4000817//SUPPRESSOR PROTEIN SRP40.//1.3e-05:255:21//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32583

20 (BAKER'S YEAST).//P3 F-NT2RP4000833

F-NT2RP4000837/MALE SPECIFIC SPERM PROTEIN MSTS4DB.//0.18:38:44//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643

F-NT2RP4000839//TRANSCRIPTION INITIATION FACTOR TFIID 90 KD SUBUNIT (TAFII-90),//0.026:38:44// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P38129

F-NTZRP4000855//AMINOPEPTIDASE B (EC 3.4.11.6) (ARGINYL AMINOPEPTIDASE) (ARGININE AMI-NOPEPTIDASE) (CYTOSOL AMINOPEPTIDASE IV) (AP-B).//2.8e-64:229:53//RATTUS NORVEGICUS (RAT).// 009175

F-NT2RP4000865//ZINC FINGER PROTEIN ZFP-36 (FRAGMENT).//3.6e-84:174:54//HOMO SAPIENS (HU-MAN).//P16415

F-NT2RP4000878//MYELOID UPREGULATED PROTEIN.//8.2e-88:227:74//MUS MUSCULUS (MOUSE).//
035682

F-NT2RP4000879//UBIQUITIN-ACTIVATING ENZYME E1 (A1S9 PROTEIN).//9.1e-55:268:43//HOMO SAPIENS (HUMAN).//P22314

35 F-NT2RP4000907//BDNF / NT-3 GROWTH FACTORS RECEPTOR PRECURSOR (EC 2.7.1.112) (TRKB TYRO-SINE KINASE) (GP145-TRKB) (TRK-B).//5.4e-10:220:25//HOMO SAPIENS (HUMAN).//Q16620

F-NT2RP4000915//60S ACIDIC RIBOSOMAL PROTEIN P2 (FRAGMENT).//0.46:23:60//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P51407

F-NT2RP4000918//METHYL-ACCEPTING CHEMOTAXIS PROTEIN TLPB.//0.00010:148:32//BACILLUS SUBTI-LIS.//P39217

F-NT2RP4000925//FIBROMODULIN PRECURSOR (FM) (COLLAGEN-BINDING 59 KD PROTEIN).//3.5e-27: 220:36//HOMO SAPIENS (HUMAN).//Q06828

F-NT2RP4000927//TRANS-ACTING TRANSCRIPTIONAL PROTEIN ICPO (P135 PROTEIN) (IER 2.9/ER2.6).// 0.64.75:37//BOVINE HERPESVIRUS TYPE 1 (STRAIN JURA).//P29128

45 F-NT2RP4000928//PHOSPHATIDATE CYTIDYLYLTRANSFERASE (EC 2.7.7.41) (CDP-DIGLYCERIDE SYN-THETASE) (CDP-DIGLYCERIDE PYROPHOSPHORYLASE) (CDP-DIACYLGLYCEROL SYNTHASE) (CDS) (CTP-PHOSPHATIDATE CYTIDYLYLTRANSFERASE) (CDP-DAG SYNTHASE) //3.1e-104:263:66//HOMO SA-PIENS (HUMAN) //029203

F-NT2RP4000929//HYPOTHETICAL 22.2 KD PROTEIN IN NSR1-TIF4631 INTERGENIC REGION.//0.93:107: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53288

F-NT2RP4000955//PUTATIVE CUTICLE COLLAGEN F09G8.6.//2.0e-05:102:37//CAENORHABDITIS ELE-GANS.//P34391

F-NT2RP4000973//HYPOTHETICAL 48.6 KD PROTEIN IN BET1-PAN1 INTERGENIC REGION.//2.3e-17:78:56// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40564

F-NT2RP4000975//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.0041:142:33//HOMO SAPIENS (HUMAN).//P10162

F-NT2RP4000979//HYPOTHETICAL 14.5 KD PROTEIN.//0.77:106:33//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20517

- F-NT2RP4000984//HYPOTHETICAL 124.8 KD PROTEIN C29E4.4 IN CHROMOSOME III.//0.90:94:25// CAENORHARDITIS FLEGANS //P34343
- F-NT2RP4000989//ANTHOPLEURIN B (TOXIN AP-B).//0.76:41:41//ANTHOPLEURA XANTHOGRAMMICA (GI-ANT GREEN SEA ANEMONE).//P01531
- F-NT2RP4000996//PROTEIN Q300.//0.00024:41:53//MUS MUSCULUS (MOUSE).//Q02722

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- F-NT2RP4009997/JDNA-DIRECTED RNA POLYMERASE I 135 KD POLYPEPTIDE (EC 2.7.7.6) (RNA POLYMER-ASE I SUBUNIT 2) (RPA135) (RNA POLYMERASE I 127 KD SUBUNIT).//8.76-115:261:82//RATTUS NORVEGI-CILS (RAT) //054888
- F-NT2RP4001004//EC PROTEIN HOMOLOG 2 (FRAGMENT).//0.50:61:34//ARABIDOPSIS THALIANA (MOUSE-FAR CRESS).//Q42377
  - F-NT2RP4001006//HYPOTHETICAL 43.5 KD PROTEIN IN COTD-KDUD INTERGENIC REGION PRECUR-SOR.//0.010:152:29//BACILLUS SUBTILIS.//P50840
  - F-NT2RP4001010/IGLUCOAMYLASE S1/S2 PRECURSOR (EC 3.2.1.3) (GLUCAN 1.4-ALPHA- GLUCOSI-DASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//9.9e-05:247.25//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//PO8640
  - F-NT2RP4001029//PROTEIN GRAINY-HEAD (DNA-BINDING PROTEIN ELF-1) (ELEMENT I-BINDING ACTIV-ITY) (TRANSCRIPTION FACTOR NTF-1).//1.1e-14:175:31//DROSOPHILA MELANOGASTER (FRUIT FLY).// B10025
  - F-NT2RP4001041//PROBABLE LEUCYL-TRNA SYNTHETASE (EC 6.1.1.4) (LEUCINE--TRNA LIGASE) (LEURS).//1.5e-74:272:55//CAENORHABDITIS ELEGANS.//Q09996
  - F-NT2RP4001057//HYPOTHETICAL 62.2 KD PROTEIN ZK652.6 IN CHROMOSOME III.//0.0064:76:38// CAENORHABDITIS ELEGANS.//P34664
    - F-NT2RP4001064//DUALIN.//2.5e-24:199:38//GALLUS GALLUS (CHICKEN).//Q90830
  - F-NT2RP4001078//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII-135) (TAFII-130) (TAFII-13
- F-NT2RP4001079//CALCIUM-TRANSPORTING ATPASE 1 (EC 3.6.1.38) (GOLGI CA2+-ATPASE) //1.5e-22:242: 31/SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P13586
  - F-NT2RP4001080//POLYPYRIMIDINE TRACT-BINDING PROTEIN (PTB) (HETEROGENEOUS NUCLEAR RI-BONUCLEOPROTEIN I) (HNRNP I).//1.7e-82:178:69//SUS SCROFA (PIG).//Q29099
- 30 F-NT2RP4001086//LEUCINE-RICH ACIDIC NUCLEAR PROTEIN.//0.00039:141:26//RATTUS NORVEGICUS (RAT).//P49911
  - F-NT2RP4001095//DOUBLE-STRANDED RNA-SPECIFIC EDITASE 1 (EC 3.5.-.) (DSRNA ADENOSINE DEAM-INASE) (RNA EDITING ENZYME 1).//9.96-07:79:43//HOMO SAPIENS (HUMAN).//P78563
  - F-NT2RP4001100//HYPOTHETICAL 74.0 KD PROTEIN IN CAJ1-HOM3 INTERGENIC REGION.//4.4e-16:207: 35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40032
    - F-NT2RP4001117//PROTEIN TRANSPORT PROTEIN SEC61 ALPHA SUBUNIT.//8.1e-115:224:99//RATTUS NORVEGICUS (RAT).//P38378
    - F-NT2RP4001122//TIPD PROTEIN://7.5e-11:129:31//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//O15736 F-NT2RP4001126//TRICHOHYALIN.//1.4e-19:257:28//OVIS ARIES (SHEEP).//P22793
- 40 F-NT2RP4001138//PUTATIVE F420-DEPENDENT NADP REDUCTASE (EC 1.-.-).//0.00010:204:25//METH-ANOCOCCUS JANNASCHII.//Q58896
  - F-NT2RP4001143//HYPOTHETICAL 52.9 KD PROTEIN IN SAP155-YMR31 INTERGENIC REGION.//4.5e-34: 168:44//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43616
- F-NT2RP4001148//SOF1 PROTEIN.//2.4e-41:158:41//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  45 P33750
  - F-NT2RP4001149//SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185).//1.3e-08:106:41//VOLVOX CART-ERI.//P21997
  - F-NT2RP4001150//NG-CAM RELATED CELL ADHESION MOLECULE PRECURSOR (NR-CAM) (BRAVO).// 3.6e-24:194:32//GALLUS GALLUS (CHICKEN).//P35331
  - F-NT2RP4001159//MEROZOITE SURFACE ANTIGEN 2 PRECURSOR (MSA-2).//0.0056:117:25//PLASMODI-UM FALCIPARUM (ISOLATE K1 / THAILAND).//Q03643
    - F-NT2RP4001174//NON-GREEN PLASTID TRIOSE PHOSPHATE TRANSLOCATOR PRECURSOR (CTPT).//
      5.96-24:184:34//BRASSICA OLERACEA (CAULIFLOWER).//P52178
  - F-NT2RP4001206//MEROZOITE SURFACE ANTIGEN 2 PRECURSOR (MSA-2).//0.0029:117:26//PLASMODI-UM FALCIPARUM (ISOLATE K1 / THAILAND) //Q03643
- 55 UM FALCIPARUM (ISOLATE K1 / THAILAND).//Q03643 F-NT2RP4001207//CHROMOSOME SEGREGATION PROTEIN CSE1.//1.0e-07:144:28//SACCHAROMYCES
  - CEREVISIAE (BAKER'S YEAST).//P33307 F-NT2RP4001210//DERMORPHIN 1 PRECURSOR [CONTAINS: DELTORPHIN (DERMENKEPHALIN); DER-

- MORPHINI.//0.019:130:30//PHYLLOMEDUSA SAUVAGEI (SAUVAGE'S LEAF FROG).//P05422
- F-NT2RP4001213//ZINC FINGER PROTEIN 177.//3.2e-28:176:39//HOMO SAPIENS (HUMAN).//Q13360
- F-NT2RP4001219//DISULFIDE ISOMERASE MPD1 PRECURSOR (EC 5.3.4.1).//2.4e-13:108:37//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//Q12404
- 5 F-NT2RP4001228//RING CANAL PROTEIN (KELCH PROTEIN).//2.7e-56:242:40//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
  - F-NT2RP4001235//REGULATORY PROTEIN E2.//0.0080:100:38//HUMAN PAPILLOMAVIRUS TYPE 25.// P36787
  - F-NT2RP4001256//CUTICLE COLLAGEN 1.//0.014:104:31//CAENORHABDITIS ELEGANS.//P08124 F-NT2RP4001258//BACTERIOCIN MICROCIN B17 PRECURSOR (MCB17).//0.00077:16:68//ESCHERICHIA COLI.//P05834
    - F-NT2RP4001274//HISTONE H1.M6.1.//0.98:65:35//TRYPANOSOMA CRUZI.//P40273

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- F-NT2RP4001276//ELAV PROTEIN.//0.00054:134:33//DROSOPHILA VIRILIS (FRUIT FLY).//P23241
- F-NT2RP4001313//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).//
  15 0.014;71;35//NICOTIANA TABACUM (COMMON TOBACCO).//P13983
  - F-NT2RP4001315//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS9.//2.3e-12:190:27//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P54787
    - F-NT2RP4001336//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//0.0037:108:31//PODOSPORA AN-SERINA //000808
- F-NT2RP4001339//HYPOTHETICAL PROTEIN MJ0810.//1.2e-09:150:34//METHANOCOCCUS JANNASCHII.// 058220
- F-NT2RP4001343//HYPOTHETICAL 85.2 KD PROTEIN F52C9.3 IN CHROMOSOME III.//1.4e-18:244:27// CAFNORHABDITIS FI FGANS //0.10123
- CAENORHABDITIS ELEGANS.//Q10123 F-NT2RP4001345//PHOSPHATIOYLCHOLINE-STEROL ACYLTRANSFERASE PRECURSOR (EC 2.3.1.43)
- 26 (LECITHIN-CHOLESTEROL ACYLTRANSFERASE) (PHOSPHOLIPID-CHOLESTEROL ACYLTRANSFERASE) (FRAGMENT)//4.0e-49:212:50/i/GALLUS GALLUS (CHICKEN)//P53760 F-NT2RP4001351/IMTRACELLULAR PROTEIN TRANSPORT PROTEIN USO1.//5.7e-11:229:26//SACCHARO-
  - MYCES CEREVISIAE (BAKER'S YEAST).//P25386
- F-NT2RP4001353//RAS SUPPRESSOR PROTEIN 1 (RSU-1) (RSP-1 PROTEIN) (RSP-1).//0.00088:84:28//HO-MO SAPIENS (HUMAN).//Q15404
  - F-NT2RP4001372//IRREGULAR CHIASM C-ROUGHEST PROTEIN PRECURSOR (IRREC PROTEIN).//1.0e-22:222:30//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q08180
    - F-NT2RP4001373//OV-17 ANTIGEN PRECURSOR (IMMUNODOMINANT HYPODERMAL ANTIGEN).//0.51:92: 26//ONCHOCERCA VOLVULUS.//P36991
- F-NT2RP4001375//NON-RECEPTOR TYROSINE KINASE SPORE LYSIS A (EC 2.7.1.112) (TYROSINE- PRO-TEIN KINASE 1).//3.5e-13:146:35//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P18160
  - F-NT2RP4001379//HYPOTHETICAL 64.2 KD PROTEIN IN SLT2-PUT2 INTERGENIC REGION.//1.2e-14:207: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38767
- F-NT2RP4001389//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (RPB1) (FRAG-49 MENT) //0.073:112:33//CRICETULUS GRISEUS (CHINESE HAMSTER) //P11414
- F-NT2RP4001407//CENTROMERIC PROTEIN E (CENP-E PROTEIN).//0.0019:233:24//HOMO SAPIENS (HU-
  - F-NT2RP4001414//SEPTIN 2 HOMOLOG (FRAGMENT).//6.2e-89:195:81//HOMO SAPIENS (HUMAN).//014141 F-NT2RP4001433//ZINC FINGER PROTEIN 43 (ZINC PROTEIN HTF6).//1.5e-85:216:56//HOMO SAPIENS (HUMAN).//P28160
    - F-NT2RP4001442//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6) (VERSION 1).// 0.012:107:35//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P18616
    - F-NT2RP4001447//60S ACIDIC RIBOSOMAL PROTEIN P2 (EL12).//0.0046:69:33//ARTEMIA SALINA (BRINE SHRIMP).//P02399
  - F-NT2RP4001474//CBP3 PROTEIN PRECURSOR.//0.0011:111:29//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P21560
    - F-NT2RP4001483//2-OXOGLUTARATE DEHYDROGENASE E1 COMPONENT PRECURSOR (EC 1.2.4.2) (AL-PHA-KETOGLUTARATE DEHYDROGENASE).//6.2e-60:146:61//HOMO SAPIENS (HUMAN).//Q02218
- F-NT2RP4001498/HYPOTHETICAL 72.5 KD PROTEIN C2F7.10 IN CHROMOSOME I.//2.3e-24:137:37// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//009701
- F-NT2RP4001502//HYPOTHETICAL 24.7 KD PROTEIN IN POM152-REC114 INTERGENIC REGION.//6.0e-22: 148:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40206
  - F-NT2RP4001507//CUTICLE COLLAGEN 40.//0.00029:166:31//CAENORHABDITIS ELEGANS.//P34804

- F-NT2RP4001524//LACTOCOCCIN A IMMUNITY PROTEIN.//0.74:96:30//LACTOCOCCUS LACTIS (SUBSP. LACTIS) (STREPTOCOCCUS LACTIS), AND LACTOCOCCUS LACTIS (SUBSP. CREMORIS) (STREPTOCOC-CUS CREMORIS)//Q00561
- F-NT2RP4001529//PROTEIN GRAINY-HEAD (DNA-BINDING PROTEIN ELF-1) (ELEMENT I-BINDING ACTIV-ITY) (TRANSCRIPTION FACTOR NTF-1).//2.8e-08:79:41//DROSOPHILA MELANOGASTER (FRUIT FLY).// B1092
  - F-NT2RP4001547//HYPOTHETICAL 45.0 KD PROTEIN IN NOT1/CDC39-HMR INTERGENIC REGION.//5.4e-34:88:46//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P25656
  - F-NT2RP4001551//CELL DIVISION CONTROL PROTEIN 68.//1.5e-18:243:30//SACCHAROMYCES CEREVI-
- SIAE (BAKER'S YEAST),//P32558
  F-NT2RP4001555//PUTATIVE ENDONUCLEASE VIII (EC 3.2:--),//0.00030:158:24//MYCOBACTERIUM TU-BERCULOSIS,//P98902

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- F-NT2RP4001567/IMPORTIN ALPHA-1 SUBUNIT (KARYOPHERIN ALPHA-1 SUBUNIT).//0.00013:147:29// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P52170
- 15 F-NT2RP4001568//HYPOTHETICAL PROTEIN KIAA0041 (FRAGMENT).//8.0e-22:119:42//HOMO SAPIENS (HUMAN).//Q15057
  - F-NT2RP4001571//NEUROMODULIN (AXONAL MEMBRANE PROTEIN GAP-43) (PP46) (B-50) (PROTEIN F1) (CALMODULIN-BINDING PROTEIN P-57).//0.012:167:28//BOS TAURUS (BOVINE).//P06836
    - F-NT2RP4001574//COATOMER GAMMA SUBUNIT (GAMMA-COAT PROTEIN) (GAMMA-COP).//6.8e-115:208: 98//BOS TAURUS (BOVINE).//P53620
      - F-NT2RP4001575//M-RELATED PROTEIN PRECURSOR.//0.22:184:25//STREPTOCOCCUS PYOGENES.// P16946
    - F-NT2RP4001592/ISOLEUCYL-TRNA SYNTHETASE (EC 6.1.1.5) (ISOLEUCINE--TRNA LIGASE) (ILERS).//
      7.4e-45;229;39//SYNECHOCYSTIS SP. (STRAIN PCC 6803).//P73505
- 25 F-NT2RP4001610//APOLIPOPROTEIN C-III PRECURSOR (APO-CIII).//0.41:74:28//SUS SCROFA (PIG).// P27917
  - F-NT2RP4001614//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//1.0:29:37//HOMO SAPIENS (HUMAN).// P02811
  - F-NT2RP4001634//MYOSIN HEAVY CHAIN, PERINATAL SKELETAL MUSCLE (FRAGMENT).//0.16:233:23// RATTUS NORVEGICUS (RAT).//P04462
    - F-NT2RP4001638//DNA REPAIR/TRANSCRIPTION PROTEIN MET18/34MS19.//4.2e-21:249:30//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P40469
      - F-NT2RP4001644//MYOSIN LIGHT CHAIN KINASE (EC 2.7.1.117) (MLCK).//4.5e-18:111:44//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P25323
- 35 F-NT2RP4001656//HYPOTHETICAL 108.5 KD PROTEIN R06F6.2 IN CHROMOSOME II.//3.4e-13:175:32// CAENORHABDITIS ELEGANS.//Q09600
  - F-NT2RP4001677//HYPOTHETICAL 73.6 KD PROTEIN CY49.21.//0.065:66:43//MYCOBACTERIUM TUBERCU-LOSIS.//Q10690
- F-NT2RP4001679//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!//1.3e-36:103:72//HOMO SAPIENS (HU-40 MAN),//P39194
  - F-NT2RP4001696//PHOTOSYSTEM II REACTION CENTRE J PROTEIN.//0.93:37:37//CHLORELLA VUL-GARIS.//P56338
  - F-NT2RP4001725//GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT.//4.3e-11:128:32// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10282
- 45 F-NT2RP4001730//UDP-GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (EC 2.4.1.-) (DUGT).//4.1e-22:201:27//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q09332
  - F-NT2RP4001739/HOMEOBOX PROTEIN HOX-A10 (HOX-1H) (HOX-1.8) (PL)//1.0:67:34//HOMO SAPIENS (HUMAN).//P31280

    F-NT2RP4001753//ZINC FINGER PROTEIN 10 /ZINC FINGER PROTEIN KOX1) (FRAGMENT).//1 2e-19:72-62//
  - F-NT2RP4001753//ZINC FINGER PROTEIN 10 (ZINC FINGER PROTEIN KOX1) (FRAGMENT).//1.2e-19:72:62/ HOMO SAPIENS (HUMAN).//P21506
  - F-NT2RP4001760//BREAKPOINT CLUSTER REGION PROTEIN.//1.8e-13:179:28//HOMO SAPIENS (HU-MAN).//P11274
    - F-NT2RP4001790//ZINC FINGER PROTEIN 38 (ZFP-38) (CTFIN51) (TRANSCRIPTION FACTOR RU49).//7.9e-38:147:49//MUS MUSCULUS (MOUSE).//Q07231
- F-NT2RP4001803//CUTICLE COLLAGEN 12 PRECURSOR.//0.40:48:39//CAENORHABDITIS ELEGANS.// P20630
  - F-NT2RP4001822//NOVEL ANTIGEN 2 (NAG-2).//2.7e-27:173:36//HOMO SAPIENS (HUMAN).//O14817 F-NT2RP4001823//PUTATIVE CUTICLE COLLAGEN F09G8.6.//3.3e-16:152:42//CAENORHABDITIS ELE-

GANS //P34391

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E-NT2RP4001828//HOLIN //0 99:33:36//BACTERIOPHAGE HP1 //P51727

F-NT2RP4001838//METASTASIS-ASSOCIATED PROTEIN MTA1.//1.2e-07:95:31//HOMO SAPIENS (HUMAN).//

- F-NT2RP4001841//INTESTINAL MUCIN-LIKE PROTEIN (MLP) (FRAGMENT).//0.94:141:22//RATTUS NOR-VEGICUS (RAT).//P98089
  - F-NTZRP4001849//SH3-BINDING PROTEIN 3BP-1//5.6e-52:276:45//MUS MUSCULUS (MOUSE).//P55194 F-NTZRP4001861//HYPOTHETICAL 10.6 KD PROTEIN IN GALE-PEPT INTERGENIC REGION.//0.92:39:51// BACILLUS SUBTILIS.//P55185
- 10 F-NT2RP4001889//HYPOTHETICAL BHLF1 PROTEIN.//0.32:97:31//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181
  - F-NT2RP4001893/Y-54-DEPENDENT RIBONUCLEASE (EC 3.1.26-) (2-54-DEPENDENT RNAASE) (RNASE L) (RIBONUCLEASE 4) (FRAGMENT) //3.6e-07:124:29/MUS MUSCULUS (MOUSE) //Q05921 F-NT2RP4001896//HYPOTHETICAL 39.4 KD TRP-ASP REPEATS CONTAINING PROTEIN IN PMT6-PCT1
- 15 INTERGENIC REGION J/3.9e-10:210:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P42935 F-NT2RP4001901//ACROSIN PRECURSOR (EC 3.4.21.10).//2.4e-07:53:45//ORYCTOLAGUS CUNICULUS (RABBIT).//P40038
  - F-NT2RP4001927//MICROTUBULE-ASSOCIATED PROTEIN YTM1.//3.1e-19:170:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q12024
- F-NT2RP4001988/ZINC FINGER PROTEIN MOK-2./1/ 3e-28-72:50/MUS MUSCULUS (MOUSE) //P24399 F-NT2RP4001946//PROTEIN-L-ISOASPARTATE O-METHYLTRANSFERASE (EC 2.1.1.77) (PROTEIN- BETA-ASPARTATE METHYLTRANSFERASE) (PIMT) (PROTEIN L-ISOASPARTYL METHYLTRANSFERASE) (L-ISO-ASPARTYL PROTEIN CARBOXYL METHYLTRANSFERASE)/4-8e-14:183:30/TRITICUM AESTIVUM (WHEAT)/G/43290
- 25 F-NT2RP4001950//HYPOTHETICAL PROTEIN ORF-1137.//3.7e-07:115:29//MUS MUSCULUS (MOUSE).// P11260 F-NT2RP4001953
  - F-NT2RP4001936 F-NT2RP4001966//WALL-ASSOCIATED PROTEIN PRECURSOR.//0.13:151:27//BACILLUS SUBTILIS.//
- 30 F-NT2RP4001975//FIBRIL-FORMING COLLAGEN ALPHA CHAIN.//0.00031:190:31//RIFTIA PACHYPTILA (TUBE WORM).//P30754
- F-NT2RP4002018//RING CANAL PROTEIN (KELCH PROTEIN).//3.5e-18:185:29//DROSOPHILA MELA-NOGASTER (FRUIT FLY)./Q04652
  - F-NT2RP4002047//GTP-BINDING PROTEIN GUF1 (GTPASE GUF1).//4.0e-49:158:65//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P46943
  - F-NT2RP4002052//HYPOTHETICAL 54.3 KD PROTEIN C23D3.03C IN CHROMOSOME I.//0.0047:148:27// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09844
    - F-NT2RP4002058//PUTATIVE PRE-MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE F56D2.6.// 0.057:66:30//CAENORHABDITIS ELEGANS.//Q20875
- 40 F-NT2RP4002071//VERY HYPOTHETICAL 13.2 KD PROTEIN CY251.09.//0.94:45:46//MYCOBACTERIUM TU-BERCULOSIS.//Q10888
  - F-NT2RP4002075/TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN).//0.44:36:38//HUMAN IMMU-NODEFICIENCY VIRUS TYPE 1 (NDK ISOLATE) (HIV-1).//P18804
- F-NT2RP4002078/ZINC FINGER PROTEIN 91 (ZINC FINGER PROTEIN HTF10) (HPF7).//2.6e-19.46:76/HO45 MO SAPIENS (HUMAN)./005481
  F-NT2RP4002081/MHO-CLASS II REGULATORY FACTOR RFX1 (RFX) (ENHANCER FACTOR C) (EF-C).//
  - 2.8e-05:196:31//HOMO SAPIENS (HUMAN).//P22670
    F-NT2RP4002083//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//0.0064:29:55//OWENIA FUSI-
  - F-N12RP4002083/HTPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//0.0064:29:55//OWENIA FUSI FORMIS.//P21260
- 59 F-NT2RP4002408//PROTEIN KINASE CEK1 (EC 2.7.1.-)//1.1e-37:159:53//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST)//P38938 F-NT2RP4002791/30S RIBOSOMAL PROTEIN \$20.//1.0:73:28//HELICOBACTER PYLORI (CAMPYLO-
  - BACTER PYLORI).//P56027
  - F-NT2RP4002888//HYPOTHETICAL PROTEIN TP0352.//0.98:52:26//TREPONEMA PALLIDUM.//083371
- 55 F-NT2RP4002905//G2/MITOTIC-SPECIFIC CYCLIN S13-7 (B-LIKE CYCLIN) (FRAGMENT).//5.9e-05:138:27// GLYCINE MAX (SOYBEAN).//P25012
  - F-NT2RP5003459/HOMEOBOX PROTEIN HOX-A3 (HOX-1.5) (MO-10).//0.027:40:40//MUS MUSCULUS (MOUSE).//P02831

- F-NT2RP5003461//HYPOTHETICAL PROTEIN C22F3.14C IN CHROMOSOME I (FRAGMENT).//1.1e-12:142: 35//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09779
- F-NT2RP5003477//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//5.3e-13:215:28//PODOSPORA AN-SERINA //Q00808
- 5 F-NT2RP5003492//GLUCOAMYLASE S1/52 PRECURSOR (EC 3.2.1.3) (GLUCAN 1,4-ALPHA- GLUCOSI-DASE) (1,4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//0.0055:144:27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)./P08640
  - F-NT2RP5003500//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//9.0e-05:103:38//MUS MUSCULUS (MOUSE).//P05142
- 10 F-NT2RP5003506//MALE SPECIFIC SPERM PROTEIN MST87F.//0.53:21:38//DROSOPHILA MELANOGASTER (FRUIT FLY).//P08175
  - F-NT2RP5003512//HYPOTHETICAL PROTEIN IN CYCB 3'REGION PRECURSOR (ORF2) (FRAGMENT).//0.92: 49:32//PARACOCCUS DENITRIFICANS.//P29969
- F-NT2RP5003522//NADPH-CYTOCHROME P450 REDUCTASE (EC 1.6.2.4) (CPR).//2.7e-18:165:39//PHASE-OLUS AUREUS (MUNG BEAN) (VIGNA RADIATA).//P37116
  - F-NT2RP5003524//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//6.0e-08:125:41//RATTUS NORVEGICUS (RAT).//P02454
    - F-NT2RP5003534//ATP SYNTHASE, SUBUNIT F (EC 3.6.1.34).//0.88:37:45//HALOBACTERIUM VOLCANII (HALOFERAX VOLCANII).//Q48331
  - F-OVARC1000001//GAR22 PROTEIN.//1.9e-05:41:58//HOMO SAPIENS (HUMAN).//Q99501
    - F-OVARC1000004//70 KD EXOCYST COMPLEX PROTEIN.//3.7e-08:186:25//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P19658
    - F-OVARC1000006//HISTONE H2A.1.//4.7e-55:117:98//RATTUS NORVEGICUS (RAT).//P02262
- F-OVARC1000013//WD-REPEAT PROTEIN POP1.//0.00022:126:28//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST) //P87080
- F-OVARC1000014//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//2.3e-05:220:30//GALLUS GALLUS (CHICKEN).//P02457
  - F-OVARC1000017//CUTICLE COLLAGEN DPY-13.//2.6e-05:97:30//CAENORHABDITIS ELEGANS.//P17657 F-OVARC1000035
- 30 F-OVARC1000058//RAS-RELATED PROTEIN RABC.//0.00015:110:24//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P34143
  - F-OVARC1000060//EXTRACELLULAR RIBONUCLEASE LE PRECURSOR (EC 3.1.27.1) (RNASE LE).//6.8e-09: 60:45//LYCOPERSICON ESCULENTUM (TOMATO).//P80022
  - F-OVARC1000068//CYTOTOXIN 4 (CARDIOTOXIN V-II-4),//1.0:27:44//NAJA MOSSAMBICA (MOZAMBIQUE COBRA),//P01452
    - F-OVARC1000071//NUCLEAR TRANSPORT FACTOR 2 (NTF-2) (PLACENTAL PROTEIN 15) (PP15).//5.2e-06: 115:29//HOMO SAPIENS (HUMAN), AND RATTUS NORVEGICUS (RAT).//P13662 F-OVARC1000085
    - F-OVARC1000087//HISTONE MACRO-H2A.1.//1.2e-13:174:26//RATTUS NORVEGICUS (RAT).//Q02874
- 40 F-OVARC1000091//OCTAPEPTIDE-REPEAT PROTEIN T2.//0.0013:137:32//MUS MUSCULUS (MOUSE).// Q06666
  - F-OVARC1000092//MITOCHONDRIAL RIBOSOMAL PROTEIN S7.//0.97:46:39//ACANTHAMOEBA CASTELLA-NII (AMOEBA).//P46756
- F-OVARC1000106/HYPOTHETICAL 141.5 KD PROTEIN IN YPT53-RHO2 INTERGENIC REGION.//0.0012:165: 29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53935
  - F-OVARC1000109//PROLINE RICH 33 KD EXTENSIN-RELATED PROTEIN PRECURSOR (FRAGMENT).//0.18: 35:34//DAUCUS CAROTA (CARROT).//P06600
  - F-OVARC1000113//HYPOTHETICAL PROTEIN C18.//1.0:26:26//SWINEPOX VIRUS (STRAIN KASZA) (SPV).// P32217
- 50 F-OVARC1000114//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.6e-28:57:63//HOMO SAPIENS (HUMAN).// P39194
  - F-OVARC1000133

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- F-OVARC1000139//UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 4 (EC 3.1.2.15) (UBIQUITIN THIOLESTE-
- RASE 4) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 4) (DEUBIQUITINATING ENZYME 4) (UBIQUI55 TOUS NUCLEAR PROTEIN HOMOLOG).//1.9e-09:200:29//HOMO SAPIENS (HUMAN).//Q13107
  - F-OVARC1000145//HOMEOBOX PROTEIN DLX-3.//1.0:65:30//BRACHYDANIO RERIO (ZEBRAFISH) (ZEBRA DANIO).//Q01702
    - F-OVARC1000148//HYPHAL WALL PROTEIN 1 (CELL ELONGATION PROTEIN 2).//0.12:175:29//CANDIDA AL-

- BICANS (YEAST) //P46593
- F-OVARC1000151//HYPOTHETICAL PROTEIN KIAA0161.//5.6e-20:197:30//HOMO SAPIENS (HUMAN).// P50876
- F-OVARC1000168//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.0030:77:38//HOMO SAPIENS (HUMAN).//
- F-OVARC1000191//COLANIC ACID BIOSYNTHESIS PROTEIN WCAH.//0.95:56:35//ESCHERICHIA COLI.// P32056
- F-OVARC1000198//HISTONE H1 C2 //0 96:70:25//TRYPANOSOMA CRUZI //P40268
- F-OVARC1000209//HYPOTHETICAL 20.9 KD PROTEIN IN PLB1-HXT2 INTERGENIC REGION.//2.5e-33:178:
  44//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)./Q03677
  - F-OVARC1000212//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//1.7e-05:66:46//MUS MUSCULUS (MOUSE).//P05142
  - F-OVARC1000240//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.8e-10:41:78//HOMO SAPIENS (HUMAN).// P39193
- 15 F-OVARC1000241//ENDOTHELIAL PAS DOMAIN PROTEIN 1 (EPAS-1) (HIF-1 ALPHA-LIKE FACTOR) (MHLF) (HIF-RELATED FACTOR) (HRF)://7.4e-54:177:54//MUS MUSCULUS (MOUSE)://P97481
  - F-OVARC1000288/HYPOTHETICAL 54.2 KD PROTEIN IN ERP5-ORC6 INTERGENIC REGION.//2.9e-20:115: 45//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38821
  - F-OVARC1000302//CORTICOSTEROID-BINDING GLOBÜLIN PRECURSOR (CBG) (TRANSCORTIN).//1.0:79: 25//MUS MUSCULUS (MOUSE).//Q06770
  - F-OVARC1000304//PROTEIN MOV-10.//1.6e-79:181:83//MUS MUSCULUS (MOUSE).//P23249
    - F-OVARC1000309//THREONINE SYNTHASE (EC 4.2.99.2),//6.9e-36:156:42//ASHBYA GOSSYPII (EREMOTH-ECIUM GOSSYPII).//Q00063
- F-OVARC1000321//HYPOTHETICAL 28.1 KD PROTEIN C4F8.03 IN CHROMOSOME I.//5.2e-45:159:53// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O14179
  - F-OVARC1000326//BASIC PROLINE-RICH PEPTIDE IB-1.//0.036:67-35//HOMO SAPIENS (HUMAN).//P04281 F-OVARC100035//HYPOTHEITOAL 39.3 KD PROTEIN IN GCN4-WBP1 INTERGENIC REGION.//1.2e-16:200: 27//SACCHAROMYCES CEREVISIAE (RAKER'S YEAST).//#40004
- F-OVARC1000347//HYPOTHETICAL 7.6 KD PROTEIN YCF33.//0.69:41:43//CYANOPHORA PARADOXA.//
  P48273
- F-OVARC1000384//ANTIFREEZE PEPTIDE 4 PRECURSOR.//0.98:49:34//PSEUDOPLEURONECTA AMERI-CANUS (WINTER FLOUNDER).//P02734
  - F-OVARC1000408/INTEGUMENTARY MUCIN C.1 (FIM-C.1) (FRAGMENT).//8.1e-05:115:33//XENOPUS LAE-VIS (AFRICAN CLAWED FROG)//005049
- 35 F-OVARC1000411//DVNACTIN, 150 KD ISOFORM (150 KD DYNEIN-ASSOCIATED POLYPEPTIDE) (DP-150) (DAP-150) (P150-GLUED).//0.00076:100:29//RATTUS NORVEGICUS (RAT).//P28023
  - F-OVARC1000414//HYPOTHETICAL 7.0 KD PROTEIN IN BLTR-SPOIIIC INTERGENIC REGION.//1.0:46:34//
    BACILLUS SUBTILIS //P54431
- F-OVARC1000420//COLLAGEN ALPHA 2(VIII) CHAIN (ENDOTHELIAL COLLAGEN) (FRAGMENT).//0.0028:97: 49 37//HOMO SAPIENS (HUMAN).//P25067
  - F-OVARC1000427//HYPOTHETICAL 13.9 KD PROTEIN IN PRFA-SPOIIR INTERGENIC REGION.//0.70:21:47// BACILLUS SUBTILIS.//P39150
  - F-OVARC1000431

- F-OVARC1000437//TENSIN.//9.2e-42:195:52//GALLUS GALLUS (CHICKEN).//Q04205
- 45 F-OVARC1000440/PINCH PROTEIN (PARTICULARY INTERESTING NEW CYS-HIS PROTEIN).//3.4e-31:37: 97//HOMO SAPIENS (HUMAN).//P48059
  - F-OVARC1000442
    - F-OVARC1000443//CUTICLE COLLAGEN 2C (FRAGMENT).//0.0056:163:34//HAEMONCHUS CONTORTUS.// P16252
- 50 F-OVARC1000461//FIXU PROTEIN.//0.36:36:44//RHIZOBIUM LEGUMINOSARUM (BIOVAR TRIFOLII).// P42710
  - F-OVARC1000465//PROTEIN TRANSPORT PROTEIN SEC7.//2.4e-14:222:26//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P11075
- F-OVARC1000466///!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//2.3e-08:29:93//HOMO SAPIENS (HUMAN).//
  P39102
  - F-OVARC1000473//DUAL SPECIFICITY PROTEIN PHOSPHATASE 7 (EC 3.1.3.48) (EC 3.1.3.16) (DUAL SPECIFICITY PROTEIN PHOSPHATASE MKP-X) (FRAGMENT).//2.8e-06:96:36//RATTUS NORVEGICUS (RAT).// O63340

- F-OVARC1000479//PHOTOSYSTEM I REACTION CENTRE SUBUNIT X (PSI-K).//0.99:48:37//CYANIDIUM CALDARIUM (GALDIERIA SULPHURARIA).//P31567 F-OVARC1000486
- F-OVARC1000496//HYPOTHETICAL PROTEIN MJ1213.//1.0:62:32//METHANOCOCCUS JANNASCHII.// Q58610
- F-OVARC1000520//MEROZOITE SURFACE PROTEIN CMZ-8 (FRAGMENT).//0.0011:66:40//EIMERIA ACER-VUILINA //P09125
  - F-OVARC1000526//PROTEIN Q300.//1.2e-05:51:43//MUS MUSCULUS (MOUSE).//Q02722
  - F-OVARC1000533//NEURONAL PROTEIN 3.1 (P311 PROTEIN).//0.74:43:41//HOMO SAPIENS (HUMAN).// Q16612
- F-OVARC1009543//POLYPEPTIDE N-ACETYLGALACTOSAMINYLTRANSFERASE (EC 24.141) [PROTEIN-UDP ACETYLGALACTOSAMINYLTRANSFERASE] (UDP-GALNAC-POLYPEPTIDE, N- ACETYLGALACTOS-AMINYLTRANSFERASE) (GALNAC-T1)/I/2.3e-23:192:35/HOMO SAPIENS (HUMAN)//Q10472 F-OVARC1000556
  - F-OVARC1000557//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.6e-08:80:47//HOMO SAPIENS (HUMAN).//
  - F-OVARC1000564/VPX PROTEIN (X ORF PROTEIN) (VIRAL ACCESSORY PROTEIN).//0.45:32:50//HUMAN IMMUNODEFICIENCY VIRUS TYPE 2 (ISOLATE D194) (HIV-2).//P17760
    - F-OVARC1000573

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- F-OVARC1000576//BETA-DEFENSIN 1 (BNDB-1).//0.47:29:41//BOS TAURUS (BOVINE).//P46159
- F-OVARC1000578//COLLAGEN ALPHA 1(II) CHAIN (FRAGMENTS).//0.023:96:36//BOS TAURUS (BOVINE).// P02459
  - F-OVARC1000588//MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L3.//0.75:57:29//HOMO SAPIENS (HU-MAN) //P09001
- F-OVARC1000605//AUTOLYSIN PRECURSOR (EC 3.4.24.38) (GAMETE LYTIC ENZYME) (GLE).//0.91:134:28//
  CHLAMYDOMONAS REINHARDTII.//P31178
- F-OVARC1000822///!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.6e-36:100:80//HOMO SAPIENS (HU-MAN)//P39189
  - F-OVARC1000640//HYPOTHETICAL 8.5 KD PROTEIN YCF40 (ORF73).//0.96:34:38//ODONTELLA SINENSIS.// P49635
- 30 F-OVARC1000649//ANTHER-SPECIFIC PROTEIN SF18 PRECURSOR (FRAGMENT).//0.0036:64:37//HELIAN-THUS ANNUUS (COMMON SUNFLOWER)./P22357
  - F-OVARC1000661/COLLAGEN ALPHA 2(I) CHAIN (FRAGMENTS).//0.21:53:47//RATTUS NORVEGICUS (RAT).//P02466
    - F-OVARC1000678//BACTERIOCIN MICROCIN B17 PRECURSOR (MCB17).//1.0:17:58//ESCHERICHIA COLI.//
    - F-OVARC1000679//DNA-DIRECTED RNA POLYMERASE OMEGA CHAIN (EC 2.7.7.6) (TRANSCRIPTASE OMEGA CHAIN) (RNA POLYMERASE OMEGA SUBUNIT).//0.096:67:29//ESCHERICHIA COLI.//P08374
    - F-OVARC1000681//PROTEIN Q300.//0.72:16:43//MUS MUSCULUS (MOUSE).//Q02722 F-OVARC1000682//PROCESSING ALPHA-1,2-MANNOSIDASE (EC 3.2.1.-) (ALPHA-1,2-MANNOSIDASE 1B).//
- 40 7.6e-70:102:99/MUS MUSCULUS (MOUSE).//P39098 F-OVARC1000689//CADMIUM-METALLOTHIONEIN (CD-MT).//0.032:30:40//HELIX POMATIA (ROMAN SNAIL)
  - (EDIBLE SNAIL).//P33187
    F-OVARC1000700//BRAIN NEURON CYTOPLASMIC PROTEIN 2.//0.17:60:40//RATTUS NORVEGICUS (RAT).//
- P02684

  F-OVARC1000703//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.57:42:42//HOMO SAPIENS (HUMAN).//
- P02811
  - F-OVARC1000722/IN-ACETYLLACTOSAMINE SYNTHASE (EC 2.4.1.90) (N-ACETYLCLUCOSAMINE (BETA 1-8gt.4)GALACTOSYLTRANSFERASE) (EC 2.4.1.38) (LACTOSE SYNTHASE A PROTEIN (EC 2.4.1.22)) (GALACTOSYLTRANSFERASE) (GT).//1.1e-20/44.70//
- 50 F-OVARC1000730//HYPOTHETICAL 83.8 KD PROTEIN C27F2.7 IN CHROMOSOME III.//5.2e-29:224:36// CAENORHABDITIS ELECANS./Q18262 F-OVARC1000746/MATERNAL EFFECT PROTEIN STAUFEN.//6.2e-12:78:48//DROSOPHILA MELA-
  - P-0VARC1000740/MINTERNAL EFFECT PROTEIN STAUFEN.//6.28-12:76.46//DROSOFRILA MELA NOGASTER (FRUIT FLY).//P25159 F-0VARC1000769
- F-0VARC1000769
- 55 F-OVARC1000771//RAS-RELATED PROTEIN RAB-2.//1.1e-46:121:79//HOMO SAPIENS (HUMAN), AND CANIS FAMILIARIS (DOG).//P08886
  - F-OVARC1000781//HOMEOBOX PROTEIN GBX-2 (GASTRULATION AND BRAIN-SPECIFIC HOMEOBOX PROTEIN 2),//0.81:36:52//HOMO SAPIENS (HUMAN),//P52951

- F-OVARC1000787//40S RIBOSOMAL PROTEIN S14 (FRAGMENT).//0.98:37:48//SUS SCROFA (PIG).//029303 F-OVARC1000800//!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.5e-31:47:82//HOMO SAPIENS (HUMAN).// P39189
- F-OVARC1000802//HYPOTHETICAL 8.8 KD PROTEIN B0302.2 IN CHROMOSOME X.//0.16:55:40// CAENORHARDITIS FLEGANS //Q10926
  - F-OVARC1000834//SERINE/THREONINE-PROTEIN KINASE PAK-ALPHA (EC 2.7.1.-) (P68-PAK) (P21- ACTI-VATED KINASE) (ALPHA-PAK) (PROTEIN KINASE MUK2).//0.87:140:31//RATTUS NORVEGICUS (RAT).//
- F-OVARC1000846//NUCLEOLIN (PROTEIN C23).//7.0e-07:109:30//MESOCRICETUS AURATUS (GOLDEN HAMSTER).//P08199
  - F-OVARC1000850//HYPOTHETICAL 56.2 KD PROTEIN IN ERG8-UBP8 INTERGENIC REGION.//6.9e-09:180: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04991
  - F-OVARC1000862/UBIQUITIN-CONJUGATING ENZYME E2-17.5 KD (EC 6.3.2.19) (UBIQUITIN- PROTEIN IGASE) (UBIQUITIN CARRIER PROTEIN)//0.0020.74:28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)//P52490
  - F-OVARC1000876//MOB1 PROTEIN (MPS1 BINDER 1).//9.8e-39:154:55//SACCHAROMYCES CEREVISIAE
    - F-OVARC1000883//METALLOTHIONEIN-I.//0.87:38:36//CANDIDA GLABRATA (YEAST) (TORULOPSIS GLA-BRATA).//P15113
- 20 F-OVARC1000885//OXIDOREDUCTASE UCPA (EC 1,-.-).//2.8e-18:170:34//ESCHERICHIA COLI.//P37440 F-OVARC1000886//COLLAGEN ALPHA 2(I) CHAIN (FRAGMENT).//0.00033:60:45//BOS TAURUS (BOVINE).// P0/245
  - F-OVARC1000890//PROBABLE E5 PROTEIN.//0.92:7:71//HUMAN PAPILLOMAVIRUS TYPE 70.//P50774
    F-OVARC1000891//HYPOTHETICAL 8.3 KD PROTEIN (ORF5)//1.0:36:36//PARAMECIUM TETRAURELIA.//
- F-OVARC1000897/IHYPOTHETICAL 6.1 KD PROTEIN PRECURSOR (ORF87).//1.0:34:44//ORGYIA PSEU-DOTSUGATA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//010337 F-OVARC1000912//PUTATIVE CUTICLE COLLAGEN C0905-8 //4 0e-07-98:35//CAENORHABDITIS ELE-
- GANS.//Q09455
  90 F-OVARC1000915//HYPOTHETICAL PROTEIN KIAA0288 (HA6116).//1.7e-47:115:76//HOMO SAPIENS (HU-
- MAN).//P56524 F-OVARC1000924//CYTOCHROME B (EC 1.10.2.2) (FRAGMENT).//0.99:54:24//BOA CONSTRICTOR (BOA).//
  - P92848
    F-OVARC1000936//HYPOTHETICAL 7.5 KD PROTEIN IN INAA-GLPQ INTERGENIC REGION.//1.0:48:33//ES-
  - CHERICHIA COLI.//P45505 F-OVARC1000937/PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//1.0:135:31/HOMO SAPIENS (HU-MAN) //P07657
  - F-OVARC1000945//EARLY E1A 11 KD PROTEIN.//0.087:81:24//MOUSE ADENOVIRUS TYPE 1 (MAV-1).// P12533
- 40 F-OVARC1000948

P15606

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- F-OVARC1000959//HYPOTHETICAL PROTEIN MJ0933.//0.99:67:28//METHANOCOCCUS JANNASCHII.//
- F-OVARC1000960//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.8e-32:56:75//HOMO SAPIENS (HUMAN).//
- 45 F-OVARC1000964/MAMBIN (GLYCOPROTEINIB-IIA ANTAGONIST) (PLATELET AGGREGATION INHIBITOR) (DENDROASPIN).//1.0:30:36//DENDROASPIS JAMESONI KAIMOSAE (EASTERN JAMESON'S MAMBA).// PPR375
  - F-OVARC1000971
  - F-OVARC1000984//HYPOTHETICAL 52.3 KD PROTEIN IN MRPL10-ERG24 INTERGENIC REGION PRECUR-SOR.//0.093:36:47//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53832
  - F-OVARC1000996//MO25 PROTEIN.//1.9e-39:80:95//MUS MUSCULUS (MOUSE).//Q06138
    - F-OVARC1000999//BRAIN-SPECIFIC HOMEOBOX/POU DOMAIN PROTEIN 1 (BRN-1 PROTEIN).//0.00020:50: 40//HOMO SAPIENS (HUMAN).//P20264
- F-OVARC1001000//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.4e-16:43:90//HOMO SAPIENS (HUMAN).//
  P39195
- F-OVARC1001004//MALE SPECIFIC SPERM PROTEIN MST84DA.//0.95:33:42//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01642
  - F-OVARC1001010//HYPOTHETICAL PROTEIN MJ0926.//0.50:71:23//METHANOCOCCUS JANNASCHII.//

#### Q58336

F-OVARC1001011//CORTISTATIN PRECURSOR.//0.81:45:37//RATTUS NORVEGICUS (RAT).//Q62949

F-OVARC1001032//FERREDOXIN LIKE PROTEIN.//1.0:26:46//RHIZOBIUM LEGUMINOSARUM (BIOVAR PHA-SEOLI).//Q05561

F-OVARC1001034//METALLOTHIONEIN-IG (MT-1G).//0.14:9:77//HOMO SAPIENS (HUMAN).//P13640

F-OVARC1001038//NUCLEOLIN (PROTEIN C23).//3.2e-07:36:80//HOMO SAPIENS (HUMAN).//P19338
F-OVARC1001040//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.5e-18:45:60//HOMO SAPIENS (HUMAN).//

F-OVARC1001044//BIS(5'-NUCLEOSYL)-TETRAPHOSPHATASE (SYMMETRICAL) (EC 3.6.1.41) (DIADENOS-

INE TETRAPHOSPHATASE).//0.88:43:39//ESCHERICHIA COLI.//P05637 F-OVARC1001051//SERINE PROTEINASE STUBBLE (EC 3.4.21,-) (STUBBLE-STUBBLOID PROTEIN).//0.34:

117:25//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q05319
F-OVARC1001055//PRE-B CELL ENHANCING FACTOR PRECURSOR.//1.6e-33:43:97//HOMO SAPIENS (HU-

F-OVARC1001055//PRE-B CELL ENHANCING FACTOR PRECURSOR.//1.66-33:43:97//HOMO SAPIENS (HU-MAN).//P43490

15 F-OVARC1001062

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F-OVARC1001065//METHIONYL-TRNA SYNTHETASE (EC 6.1.1.10) (METHIONINE-TRNA LIGASE) (METRS).//0.79:76:39//BORRELIA BURGDORFERI (LYME DISEASE SPIROCHETE).//Q44951

F-OVARC1001068//GTP-BINDING PROTEIN ERA HOMOLOG (FRAGMENT).//5.3e-15:100:44//BRADYRHIZO-BIUM JAPONICUM.//O69162

F-OVARC1001072///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.0076:41:56//HOMO SAPIENS (HUMAN).// P39188

F-OVARC1001074//60S RIBOSOMAL PROTEIN L38.//1.0:32:40//LYCOPERSICON ESCULENTUM (TOMATO).// P46291

F-OVARC1001085//HYPOTHETICAL 126.5 KD PROTEIN C13F4.06 IN CHROMOSOME I.//0.73:135:25// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10197

F-OVARC1001092//HYPOTHETICAL 51.2 KD PROTEIN IN PET54-DIE2 INTERGENIC REGION.//5.6e-05:30: 56//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P50079

F-OVARC1001107//SHK1 KINASE-BINDING PROTEIN 1.//1.8e-08:52:51//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P78963

30 F-OVARC1001113//DIAPHANOUS PROTEIN.//1.9e-33:218:35//DROSOPHILA MELANOGASTER (FRUIT FLY).//P48608

F-OVARC1001117//GENE 7 PROTEIN.//0.68:12:50//SPIROPLASMA VIRUS 4 (SPV4).//P11339

F-OVARC1001129//30S RIBOSOMAL PROTEIN S17.//0.15:57:22//AQUIFEX AEOLICUS.//Q66439

35 F-OVARC1001154//GRANULINS PRECURSOR (ACROGRANIN),//2.3e-95:99:77//MUS MUSCULUS (MOUSE),//P28798

F-OVARC1001161//GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT.//0.17:87:34//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P49177

F-OVARC1001162

40 F-OVARC1001167//TRBD PROTEIN.//0.92:24:45//ESCHERICHIA COLI.//P41070

F-OVARC1001169//FRUCTOSE-1,6-BISPHOSPHATASE (EC 3.1.3.11) (D-FRUCTOSE-1,6-BISPHOSPHATE 1-PHOSPHOHYDROLASE) (FBPASE) (FRAGMENT).//0.82:35:40//MUS MUSCULUS (MOUSE).//P97323

F-OVARC1001170//PROLINE-RICH PEPTIDE P-B.//0.17:27:37//HOMO SAPIENS (HUMAN).//P02814

F-OVARC1001171//IIII ALU SUBFAMILY J WARNING ENTRY IIII/I0.00023:28:75//HOMO SAPIENS (HUMAN).//
45 P39188

F-OVARC1001173

F-OVARC1001176//HYPOTHETICAL BHLF1 PROTEIN.//2.7e-05:158:31//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181

F-OVARC1001180//UBIQUITIN-LIKE PROTEIN DSK2.//1.4e-12:208:25//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P48510

F-OVARC1001188//HYPOTHETICAL 27.8 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//3.3e-31: 129:51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53215

F-OVARC1001200//HYPOTHETICAL 49.0 KD PROTEIN IN NSP1-KAR2 INTERGENIC REGION.//0.018:148:26// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47057

55 F-OVARC1001232//HYPOTHETICAL PROTEIN MJ1236.//2.5e-27:141:39//METHANOCOCCUS JANNASCHII.// Q58633

F-OVARC1001240

F-OVARC1001243

- F-OVARC1001244//RING3 PROTEIN (KIAA9001).//1.7e-13:37:91//HOMO SAPIENS (HUMAN).//P25440
- F-OVARC1001261//OCTAPEPTIDE-REPEAT PROTEIN T2.//1.3e-07:109:35//MUS MUSCULUS (MOUSE).// Q06666
- F-OVARC1001268//HYPOTHETICAL 57.4 KD PROTEIN IN PILT REGION (ORF4).//0.71:43:41//PSEU-DOMONAS AFRUGINOSA //P24563
- F-OVARC1001270//HYPOTHETICAL 9.0 KD PROTEIN IN UVSW-UVSY INTERGENIC REGION.//1.0:44:29// BACTERIOPHAGE 14.//P32281
  - F-OVARC1001271//HYPOTHETICAL 104.7 KD PROTEIN F23F12.8 IN CHROMOSOME III PRECURSOR.// 0.00015:188:23//CAENORHABDITIS ELEGANS.//P46504
- 10 F-OVARC1001282
  - F-OVARC1001296//WEB1 PROTEIN (PROTEIN TRANSPORT PROTEIN SEC31),//0.022:101:31//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P38968
  - F-OVARC1001306//HYPOTHETICAL 52.9 KD SERINE-RICH PROTEIN C11G7.01 IN CHROMOSOME I.//0.023: 134;26//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//013695
- 15 F-OVARC1001329//CHLOROPLAST TRIOSE PHOSPHATE TRANSLOCATOR PRECURSOR (CTPT).//1.3e-14: 150;28//ZEA MAYS (MAIZE).//P49133
  - F-OVARC1001330

- F-OVARC1001339//RIBONUCLEOPROTEIN RB97D.//0.0013:55:38//DROSOPHILA MELANOGASTER (FRUIT FLY).//002926
- F-OVARC1001341//HYPOTHETICAL 74.0 KD PROTEIN IN CAJ1-HOM3 INTERGENIC REGION.//4.9e-17:110: 43//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40032
- F-OVARC1001342
  F-OVARC1001344//PREPROTEIN TRANSLOCASE SECE SUBUNIT.//0.99:39:23//STAPHYLOCOCCUS CAR-
- NOSUS.//P36253
  F-OVARC1001357//METALLOTHIONEIN.//0.99:28:42//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//
  OOK880
  - F-OVARC1001360/LARGE PROLINE-RICH PROTEIN BAT2 (HLA-B-ASSOCIATED TRANSCRIPT 2).//0.86:109: 31//HOMO SAPIENS (HUMAN)./P48634
- F-OVARC1001369//COLLAGEN ALPHA 2(I) CHAIN (FRAGMENT).//6.7e-05:124:36//BOS TAURUS (BOVINE).//
  P02465
- F-OVARC1001372//HYPOTHETICAL 34.5 KD PROTEIN IN CLCB-CLCD INTERGENIC REGION PRECUR-SOR.//0.75:33:48//PSEUDOMONAS PUTIDA, AND PSEUDOMONAS SP. (STRAIN B13).//Q47100
  - F-OVARC1001376//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.8e-24:96:61//HOMO SAPIENS (HUMAN).// P39188
- F-OVARC1001381//MEMBRANE-ASSOCIATED ATPASE EPSILON CHAIN (EC 3.6.1.34) (SUL-ATPASE EPSI-LON).//0.96:46:39//SULFOLOBUS ACIDOCALDARIUS.//P23039
  - F-OVARC1001391//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.00024:189:29//HOMO SAPIENS (HUMAN).//P10162
- F-OVARC1001399//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.062:18:77//HOMO SAPIENS (HUMAN).//
  P39195
  - F-OVARC1001417//HYPOTHETICAL 157.0 KD PROTEIN C38C10.5 IN CHROMOSOME III.//0.010:185:23//
    CAENORHABDITIS ELEGANS.//Q03570
  - F-OVARC1001419//A-TYPE INCLUSION PROTEIN (ATI).//0.50:135:28//CAMELPOX VIRUS (STRAIN CP-1).//
- 45 F-OVARC1001425//COLLAGEN ALPHA 1(X) CHAIN PRECURSOR.//0.43:85:40//HOMO SAPIENS (HUMAN).//
  O03692
  - F-OVARC1001436//HYPOTHETICAL 11.4 KD PROTEIN (C4 PROTEIN).//0.031:100:30//TOMATO YELLOW LEAF CURL VIRUS (STRAIN AUSTRALIA) (TYLCV).//P36283
  - F-OVARC1001442//HOMEOBOX PROTEIN HTR-A2 (FRAGMENT).//1.0:32:34//HELOBDELLA TRISERIALIS (LEECH).//P17138
  - F-OVARC1001453//METALLOTHIONEIN-III (MT-III) (GROWTH INHIBITORY FACTOR) (GIF).//0.74:19:47//MUS MUSCULUS (MOUSE).//P28184
    - F-OVARC1001476//GTP-BINDING PROTEIN GTR2.//3.0e-12:114:34//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P53290
- 55 F-OVARC1001480//COLLAGEN ALPHA 2(VI) CHAIN PRECURSOR://0.00019:134:32//MUS MUSCULUS (MOUSE).//Q02788
  - F-OVARC1001489//HYPOTHETICAL PROTEIN HI1270.//0.98:30:43//HAEMOPHILUS INFLUENZAE.//P44149 F-OVARC1001496//C-TERMINAL BINDING PROTEIN 2.//4.0e-65:132:100//HOMO SAPIENS (HUMAN).//

#### P56545

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- F-OVARC1001506//POLYCYSTIN PRECURSOR (AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE PROTEIN 1).//3.2e-70:159:94//HOMO SAPIENS (HUMAN).//P98161
- F-OVARC1001525//FIBROBLAST GROWTH FACTOR INDUCIBLE PROTEIN 14 (FIN14).//1.0:36:33//MUS MUS-CULUS (MOUSE).//Q61077
  - F-OVARC1001542//SMALL PROLINE-RICH PROTEIN 2B (SPR-2B).//0.69:57:33//HOMO SAPIENS (HUMAN).// P35325

## F-OVARC1001547

- F-OVARC1001555//NGG1-INTERACTING FACTOR 3.//7.6e-16:148:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53081
  - F-OVARC1001577/SPLICING FACTOR, ARGININE/SERINE-RICH 2 (SPLICING FACTOR SC35) (SC-35) (SC-35) (SC-35) (SC-35) (SC-35) (SPLICING COMPONENT, 35 KD) (PR264 PROTEIN)//8.86-38.934/81//GALLUS GALLUS (CHICKEN)//P30352 F-OVARC1001600//GENE 7 PROTEIN//0.80:38:39//SPIROPLASMA VIRUS SPV1-R8AZ B.//P15898
- F-OVARC1001610//DIACYLGLYCEROL CHOLINEPHOSPHOTRANSFERASE (EC 2.7.8.2) (SN-1,2- DIACYLG-LYCEROL CHOLINEPHOSPHOTRANSFERASE) (CHOPT),//1.6e-22:122:39//SACCHAROMYCES CEREVI-SIAE RIACKET'S VEASTI//#17888
  - F-OVARC1001611
  - F-OVARC1001615//HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN CHROMOSOME X.//0.30:43:34//
    CAENORHABDITIS ELEGANS.//Q11116
- F-OVARC1001668//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.0e-19:45:82//HOMO SAPIENS (HUMAN).// P39192
  - F-OVARC1001702//SOX-20 PROTEIN.//2.4e-28:71:83//HOMO SAPIENS (HUMAN).//O60248
- F-OVARC1001703//INTERFERON-INDUCED GUANYLATE-BINDING PROTEIN 1 (GUANINE NUCLEOTIDE-BINDING PROTEIN 1) (INTERFERON-GAMMA INDUCIBLE PROTEIN MAG-1).//0.00018.88:36//MUS MUSCU-LUS (MOUSE)./001514
  - F-OVARC1001711//CORNIFIN B (SMALL PROLINE-RICH PROTEIN 1B) (SPR1B) (SPR1 B).//2.7e-05:98:32// MUS MUSCULUS (MOUSE).//Q62267
    - F-OVARC1001713//ENDOZEPINE-RELATED PROTEIN PRECURSOR (MEMBRANE-ASSOCIATED DIAZEPAM BINDING INHIBITOR) (MA-DBI).//4.5e-20:46:67//BOS TAURUS (BOVINE).//P07106
- F-OVARC1001726//ALPHA-AMYLASE INHIBITOR PAIM I (PIG PANCREATIC ALPHA-AMYLASE INHIBITOR OF MICROBES 1)//0.59:23:56//STREPTOMYCES OLIVACEOVIRIDIS (STREPTOMYCES CORCHORUSI).// P089/1
  - F-OVARC1001731//TROPOMYOSIN ALPHA CHAIN, SKELETAL MUSCLE.//2.1e-75:176:87//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//001173
- 35 F-OVARC1001745//GENE 11 PROTEIN.//0.31:36:52//SPIROPLASMA VIRUS SPV1-R8A2 B.//P15902
  - F-OVARC1001762/IN-TERMINAL ACETYLTRANSFERASE 1 (EC 2.3.1.88) (AMINO-TERMINAL, ALPHA- AMI-NO, ACETYLTRANSFERASE 1).//2.8e-23:197:35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P12945
- F-OVARC1001766//FK506-BINDING NUCLEAR PROTEIN (PEPTIDVL-PROLYL CIS-TRANS ISOMERASE) (40 (PPIASE) (EC 5.2.1.8) (PROLINE ROTAMASE) (NUCLEOLAR PROLINE ISOMERASE) (FKBP-70).//2.2e-06-99: 40//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P38911
  - F-OVARC1001767/33.2 KD PROTEIN IN DIND-RPH INTERGENIC REGION (ORF X),//0.99:113:27//ES-CHERICHIA COLL/J/P23839
    F-OVARC1001768
- 45 F-OVARC1001791//HYPOTHETICAL 63.3 KD PROTEIN IN MPT5-SAE2 INTERGENIC REGION.//0.090:75:32// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P46945
  - F-OVARC1001795//HYPOTHETICAL 7.5 KD PROTEIN IN RPBA-GP46 INTERGENIC REGION.//0.81:21:38// BACTERIOPHAGE T4.//P07878
  - F-OVARC1001802//PLECTOXIN VIII (PLT-VIII) (PLTVIII).//0.41:19:36//PLECTREURYS TRISTIS (SPIDER).// P36984
    - F-OVARC1001805//60S RIBOSOMAL PROTEIN L40 (CEP52).//0.67:24:58//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P14796
      - F-OVARC1001809//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//0.23:111:31//RATTUS NORVEGICUS (RAT).//P02454
- 55 F-OVARC1001812//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.99:28:42//HALICHOERUS GRYPUS (GRAY SEAL).//P38592
  - F-OVARC1001813//HYPOTHETICAL 9.9 KD PROTEIN.//0.41:36:30//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20562

- F-OVARC1001820//HYPOTHETICAL PROTEIN ORF-1137.//0.80:58:29//MUS MUSCULUS (MOUSE).//P11260 F-OVARC1001828
- F=OVARC1001846
- F-OVARC1001861//METALLOTHIONEIN (MT).//0.18:11:54//PLEURONECTES PLATESSA (PLAICE).//P07216
- F-OVARCIO01873
  F-OVARCIO01879/HYPOTHETICAL 55.9 KD PROTEIN EEED8.6 IN CHROMOSOME II.//2.3e-05:73:31//
  - CAENORHABDITIS ELEGANS.//Q09296 F-OVARC1001880//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONE CP7) [CONTAINS: BASIC
- PEPTIDE P-FJ (FRAGMENT).//2.4e-11:203.32/HOMO SAPIENS (HUMAN).//P02812 10 F-OVARC1001883/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/2.3e-16:86:59/HOMO SAPIENS (HUMAN).// P39188
  - F-OVARC1001900/HYPOTHETICAL 105.9 KD PROTEIN F22B7.5 IN CHROMOSOME III.//0.0053:48:47// CAENORHABDITIS ELEGANS.//P34408 F-OVARC1001991
- 15 F-OVARC1001911//40S RIBOSOMAL PROTEIN \$28.//1.0:33:36//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P34789
  - F-OVARC1001916//PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR (RHO/RAC GEF) (FACIOGENITAL DYSPLASIA PROTEIN).//0.00082:114:27//HOMO SAPIENS (HUMAN).//P98174
- F-OVARC1001928//FERREDOXIN III (FDIII).//1.0:64:29//ANABAENA VARIABILIS.//P46050

  F-OVARC1001942//N-TERMINAL ACETYLTRANSFERASE 1 (EC 2.3.1.88) (AMINO-TERMINAL, ALPHA- AMI-
- F-OVARCTI01942/IN-TERMINAL ACETYLTRANSFERASE 1 (EC 2.3.1.88) (AMINO-TERMINAL, ALPHA-AMI-NO, ACETYLTRANSFERASE 1),//3.0e-07:93:37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P12945
- F-OVARC1001943//HYPOTHETICAL 62.2 KD PROTEIN ZK652.6 IN CHROMOSOME III.//1.7e-23:147:43//
  CAENORHABDITIS ELEGANS //P34664
- 25 F-OVARC1001949//ZINC FINGER PROTEIN 177.//2.0e-23;56:66//HOMO SAPIENS (HUMAN).//Q13360
  - F-OVARC1001950//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.011:57:47//HOMO SAPIENS (HUMAN).// P39188
  - F-OVARC1001987//SPERM PROTAMINE P1 (CYSTEINE-RICH PROTAMINE).//0.39:14:64//MUS MUSCULUS (MOUSE).//P02319
- 30 F-OVARC1001989/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/2.4e-13:55:72//HOMO SAPIENS (HUMAN).// P39188
  - F-OVARC1002044
    - F-OVARC1002050//UTROPHIN (DYSTROPHIN-RELATED PROTEIN 1) (DRP1) (DRP1).//3.6e-12:221:25//HOMO SAPIENS (HUMAN).//P46939
- 35 F-OVARC1002066
- F-OVARC1002082

- F-OVARC1002107//INTRACELLULAR PROTEIN TRANSPORT PROTEIN USO1.//0.99:149:24//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P25386
- F-OVARC1002112//HISTONE MACRO-H2A.1.//2.8e-64:133:98//RATTUS NORVEGICUS (RAT).//Q02874
- 40 F-OVARC1002127//60S RIBOSOMAL PROTEIN L22.//0.0023:95:35//DROSOPHILA MELANOGASTER (FRUIT FLY).//P50887
  - F-OVARC1002138/IPROBABLE 26S PROTEASE SUBUNIT YTA6 (TAT-BINDING HOMOLOG 6).//6.4e-51:198: 56//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40328 F-OVARC1002143
- 45 F-OVARC1002156//HYPOTHETICAL 27.7 KD PROTEIN IN CPT1-SPC98 INTERGENIC REGION.//0.00010:64: 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P53915
  - F-OVARC1002158/HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION.//8.2e-07:119:35// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41479
  - F-OVARC1002165//EBNA-6 NUCLEAR PROTEIN (EBNA-3C) (EBNA-4B).//0.00023:90:45//EPSTEIN-BARR VI-RUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03204
- F-OVARC1002182/JHYPOTHETICAL 46.2 KD TRP-ASP REPEATS CONTAINING PROTEIN D2013.2 IN CHRO-MOSOME II./1.3e-34:165:35//CAENORHABDITIS ELEGANS.//Q18964
- F-PLACE1000004//HYPOTHETICAL 180.2 KD PROTEIN C31A2.05C IN CHROMOSOME I.//8.8e-05:148:25// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09725
- 55 F-PLACE1000005//PROTEIN Q300.//0.30:10:100//MUS MUSCULUS (MOUSE).//Q02722
  - F-PLACE1000007/PROBABLE UBIOUTIN CARBOXYL-TERMINAL HYDROLASE R10E11.3 (EC 3.1.2.15) (UBIOUITIN THIOLESTERASE) (UBIOUITIN-SPECIFIC PROCESSING PROTEASE) (DEUBIQUITINATING ENZYME)/I.2.3a-39:134:62/I/CAENORHABDITIS ELEGANS://P34547

- F-PLACE1000014//52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS-A)) (RO(SS-A)).// 0.00036:63:39//HOMO SAPIENS (HUMAN).//P19474
- F-PLACE1000031
- F-PLACE1000040//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//4.4e-12:97:41//HOMO SAPIENS (HUMAN).//
- F-PLACE1000048//50S RIBOSOMAL PROTEIN L15 (FRAGMENT).//0.98:31:38//BACILLUS SP. (STRAIN C-125) //P38373
- F-PLACE1000050//COLLAGEN ALPHA 1(III) CHAIN.//0.00062:190:33//BOS TAURUS (BOVINE).//P04258 F-PLACE1000061//60S RIBOSOMAL PROTEIN L37A.//6.4e-19:51:86//GALLUS GALLUS (CHICKEN).//P32046
- F-PLACE1000061/60S RIBOSOMAL PROTEIN L37A.//6.4e-19:51:86//GALLUS GALLUS (CHICKEN).//P32046

  F-PLACE1000066//SSU72 PROTEIN.//2.3e-39:165:49//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
  P33538
  - F-PLACE1000078//BAD PROTEIN (BCL-2 BINDING COMPONENT 6).//1.7e-06:21:95//HOMO SAPIENS (HU-MAN).//Q92934
  - F-PLACE1000081//HOMEOBOX PROTEIN HOX-A4 (HOX-1.4) (MH-3).//0.0053:146:33//MUS MUSCULUS (MOUSE).//P06798
- 15 (MOUSE).//P06798 F-PLACE1000094
  - F-PLACE1000133//TRANSCRIPTION FACTOR BTF3 (RNA POLYMERASE B TRANSCRIPTION FACTOR 3).//
    1.8e-62:158:81//HOMO SAPIENS (HUMAN).//P20290
- F-PLACE:100142/IENOYL-COA HYDRATASE, MITOCHONDRIAL PRECURSOR (EC 4.2.1.17) (SHORT CHAIN ENOYL-COA HYDRATASE) (SCEH) (ENOYL-COA HYDRATASE 1),//9.8e-12:104:34//HOMO SAPIENS (HUMAN),//P30084
  - F-PLACE1000184//AC PROTEIN.//0.44:31:29//BACTERIOPHAGE T4.//P18924
  - F-PLACE1000185//HYPOTHETICAL GLYCINE-RICH 49.6 KD PROTEIN CY130.10C PRECURSOR.//0.11:48: 33//MYCOBACTERIUM TUBERCULOSIS.//Q10637
- F-PLACE:1000213//GLUCOAM/LASE S1/82 PRECURSOR (EC 3.2.1.3) (GLUCAN 1.4-ALPHA- GLUCOSI-DASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE)//3.4e-05:194:26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST)//P08640 F-PLACE:1000214
  - F-PLACE1000236//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR://0.027:63:34//GALLUS GALLUS (CHICKEN)://P02457
  - F-PLACE1000248/TEGUMENT PROTEIN (GENE 11 PROTEIN).//0.78:100:28/FEQUINE HERPESVIRUS TYPE 4 (STRAIN 1942) (EHV-4) (EQUINE HERPESVIRUS TYPE 1 SUBTYPE 2).//Q00039 F-PLACE1000292
    - F-PLACE1000308//EARLY NODULIN 75 (N-75) (NGM-75) (FRAGMENT).//0.049:28:42//MEDICAGO SATIVA (ALFALFA).//P11728
- 35 (ALFALFA).//P11728 F-PLACE1000332
  - F-PLACE1000347//HYPOTHETICAL PROTEIN TP0420.//0.15:24:54//TREPONEMA PALLIDUM://O83435
  - F-PLACE1000374//LYSOZYME C (EC 3.2.1.17) (1,4-BETA-N-ACETYLMURAMIDASE C).//1.0:63:25//ORYC-TOLAGUS CUNICULUS (RABBIT).//P16973
- 40 F-PLACE1000380/IMATING PROCESS PROTEIN MID2 (SERINE-RICH PROTEIN SMS1) (PROTEIN KINASE A INTERFERENCE PROTEIN)/I0.018:169.28/I/SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36027 F-PLACE1000383/IMYOTUBULARIN.//1.2e-65:215.57/IHOMO SAPIENS (HUMAN).//013496
  - F-PLACE1000401//ELASTIN PRECURSOR (TROPOELASTIN).//0.00023:145:30//MUS MUSCULUS (MOUSE).//P54320
- 45 F-PLACE1000406//54 KD NUCLEAR RNA-BINDING PROTEIN (P54(NRB)).//3.4e-27:90:63//HOMO SAPIENS (HUMAN).//Q15233
  - F-PLACE1000420//7,8-DIHYDRO-8-OXOGUANINE TRIPHOSPHATASE (EC 3.1.6.-) (8-OXO-DGTPASE).//4.7e-07:134:29//MUS MUSCULUS (MOUSE).//P53368
  - F-PLACE1000421//HYPOTHETICAL 8.8 KD PROTEIN C11D3.01C IN CHROMOSOME I.//0.48:72:27// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10080
  - F-PLACE1000424

- F-PLACE1000435
- F-PLACE1000444//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.0e-31:129:63//HOMO SAPIENS (HU-MAN) //P30105
- 55 F-PLACE1000453//PROTEIN Q300.//0.013:16:68//MUS MUSCULUS (MOUSE).//Q02722
  - F-PLACE1000481/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.14:63:36//HOMO SAPIENS (HU-MAN).//P08547
    - F-PLACE1000492//BASP1 PROTEIN.//0.17:114:28//HOMO SAPIENS (HUMAN).//P80723

#### F-PLACE1000540

F-PLACE1000547//MANNOSE-1-PHOSPHATE GUANYLTRANSFERASE (EC 2.7.7.13) (ATP-MANNOSE-1-PHOSPHATE GUANYLYLTRANSFERASE) (NDP-HEXOSE PYROPHOSPHORYLASE) //1.88-21:87:56//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST)://F41940

F-PLACE1000562//HYPOTHETICAL PROTEIN MJ0562.//1.0:35:34//METHANOCOCCUS JANNASCHII.//

F-PLACE1000564//ADRENAL SPECIFIC 30 KD PROTEIN (CLONE PG2).//0.13:66:37//HOMO SAPIENS (HU-MAN) //P15803

F-PLACE1000583//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//7.0e-45:192:47//HOMO SA-PIENS (HUMAN).//P51522

F-PLACE1000588//INTERFERON-INDUCED GUANYLATE-BINDING PROTEIN 1 (GUANINE NUCLEOTIDE-BINDING PROTEIN 1)://5.3e-63:122:88/HOMO SAPIENS (HUMAN)://P32455

F-PLACE1000596//RING CANAL PROTEIN (KELCH PROTEIN).//2.6e-12:120:38//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652

15 F-PLACE1000599//EARLY E3B 12.7 KD PROTEIN PRECURSOR.//0.83:53:32//HUMAN ADENOVIRUS TYPE 12.//P36707

#### F-PLACE1000610

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F-PLACE1000611//HYPOTHETICAL 33.6 KD PROTEIN IN MCK1-RPS19B INTERGENIC REGION.//9.4e-07:64: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P48558

F-PLACE1000636//MALE STERILITY PROTEIN 2.//3.7e-09:83:43//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//Q08891

F-PLACE1000653/PUTATIVE PHOSPHOACETYLGLUCOSAMINE MUTASE (EC 5.4.2.3) (ACETYLGLU-COSAMINE PHOSPHOMUTASE) (N-ACETYLGLUCOSAMINE-PHOSPHATE MUTASE),//1.96-30:203.41// SCHIZOSACCHAROMYCES POMBE (FISSION YEASTI//009687

F-PLACE1000656/EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).// 0.0029:75:33//NICOTIANA TABACUM (COMMON TOBACCO).//P13983

F-PLACE1000706/TRANSCRIPTION INTERMEDIARY FACTOR 1-BETA (NUCLEAR COREPRESSOR KAP-1) (KRAB-ASSOCIATED PROTEIN 1).//1.1e-38:180:42//HOMO SAPIENS (HUMAN).//Q13263

F-PLACE1000712//VERY HYPOTHETICAL 8.9 KD PROTEIN CY441.05 PRECURSOR.//0.93:49:34//MYCOBAC-TERIUM TUBERCULOSIS.//P71934

### F-PLACE1000716

F-PLACE1000748/HYPOTHETICAL 10.4 KD PROTEIN IN SPAT 3'REGION (ORF-11).//0.90:53:37//SHIGELLA FLEXNERI.//P55794

F-PLACE1000749//HYPOTHETICAL PROTEIN MG148.//0.0014:142:27//MYCOPLASMA GENITALIUM.// P47394

F-PLACE1000755//HYPOTHETICAL HELICASE K12H4.8 IN CHROMOSOME III.//1.1e-15:98:48//CAENORHAB-DITIS ELEGANS //P34529

F-PLACE1000769//VIGILIN.//0.51:60:33//GALLUS GALLUS (CHICKEN).//P81021

F-PLACE1000785//PROBABLE COLD SHOCK PROTEIN CY15C10.04.//1.0:22:45//MYCOBACTERIUM TUBER-40 CULOSIS.//006360

F-PLACE1000786//HYPOTHETICAL 30.2 KD PROTEIN ZK632.12 IN CHROMOSOME III.//2.6e-38:159:51//
CAENORHABDITIS ELEGANS.//P34657

F-PLACE1000793//VASODILATOR-STIMULATED PHOSPHOPROTEIN (VASP).//0.0097:128:30//HOMO SAPIENS (HUMAN).//P50552

45 F-PLACE1000798//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/9.4e-07:47:61//HOMO SAPIENS (HUMAN).// P39188

#### F-PLACE1000841

F-PLACE1000849//ELAV PROTEIN.//3.5e-05:140:35//DROSOPHILA VIRILIS (FRUIT FLY).//P23241

F-PLACE1000856//HYPOTHETICAL PROTEIN MJ0008.//0.95:100:23//METHANOCOCCUS JANNASCHII.// Q60319

F-PLACE1000863//PUTATIVE MITOCHONDRIAL 40S RIBOSOMAL PROTEIN YHR148W.//2.3e-46:172:54// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32899

F-PLACE1000909//ANKYRIN R (ANKYRINS 2.1 AND 2.2) (ERYTHROCYTE ANKYRIN).//0.00022:105:35//HO-MO SAPIENS (HUMAN).//P16157

55 F-PLACE1000931/KILLER TOXIN HM-1.//0.95:24:33//WILLIOPSIS MRAKII (YEAST) (HANSENULA MRAKII).//

F-PLACE1000948//SL CYTOKINE PRECURSOR (FLT3 LIGAND).//0.97:52:40//HOMO SAPIENS (HUMAN).// P49771

- F-PLACE1000972//MYOSIN ID HEAVY CHAIN.//1.9e-06:79:43//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD) //P34109
- F-PLACE1000977//HYPOTHETICAL 94.2 KD PROTEIN C38D4.5 IN CHROMOSOME III.//2.5e-23:105:41// CAENORHABDITIS ELEGANS.//P46941
- F-PLACE1000979//ZINC FINGER PROTEIN 7 (ZINC FINGER PROTEIN KOX4) (ZINC FINGER PROTEIN HF. 16).//0.91:83:30//HOMO SAPIENS (HUMAN).//P17097
  - F-PLACE1000987//HYPOTHETICAL 111.5 KD PROTEIN C22G7.02 IN CHROMOSOME I.//0.10:128:24// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09796 E-PLACE 100 1000
- 10 F-PLACE1001007//ZYXIN.//2.2e-05:135:30//GALLUS GALLUS (CHICKEN).//Q04584
  - F-PLACE1001010//BETA-1 BUNGAROTOXIN B CHAIN, MAJOR COMPONENT PRECURSOR (BUNGAROTOX-IN. B1 CHAIN).//1.0:30:40//BUNGARUS MULTICINCTUS (MANY-BANDED KRAIT).//P00987
  - F-PLACE1001015 F-PLACE1001024
- 15 F-PLACE 100 1036
  - F-PLACE1001054//HOLOTRICIN 3 PRECURSOR.//0.0044:56:39//HOLOTRICHIA DIOMPHALIA.//Q25055
  - F-PLACE1001062//SACCHAROPINE DEHYDROGENASE [NADP+, L-GLUTAMATE FORMING] (EC 1.5.1.10),// 0.0013:38:52//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P38999 F-PLACE1001076
- 20 F-PLACE1001088//EARLY NODULIN 75 (N-75) (NGM-75) (FRAGMENT).//0.95:32:50//MEDICAGO SATIVA (AL-FALFA).//P11728
  - F-PLACE1001092//HYPOTHETICAL 49.0 KD PROTEIN IN NSP1-KAR2 INTERGENIC REGION.//0.0026:81:35// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47057
- F-PLACE1001104//HYPOTHETICAL 131.5 KD PROTEIN C02F12.7 IN CHROMOSOME X.//0.00063:125:32// 25 CAENORHABDITIS ELEGANS.//Q11102
- F-PLACE1001118//ZINC FINGER PROTEIN MLZ-4 (ZINC FINGER PROTEIN 46),//2.6e-77:209:63//MUS MUS-CULUS (MOUSE),//Q03309
  - F-PLACE1001136//ALPHA-N-ACETYLGALACTOSAMINIDASE PRECURSOR (EC 3.2.1.49) (ALPHA- GALAC-TOSIDASE B).//0.99:107:30//HOMO SAPIENS (HUMAN).//P17050
- 30 F-PLACE1001168
  - F-PLACE1001171//RETROVIRUS-RELATED POL POLYPROTEIN (FRAGMENT).//0.00012:37:59//HOMO SAPI-ENS (HUMAN) //P12895
  - F-PLACE1001185//HYPOTHETICAL 56.6 KD PROTEIN IN URE2-SSU72 INTERGENIC REGION.//3.6e-12:88: 36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53867
- 35 F-PLACE1001238
  - F-PLACE1001241//METALLOTHIONEIN B (MTB) (FRAGMENT).//0.13:30:53//COLINUS VIRGINIANUS (BOB-WHITE QUAIL) (COMMON BOBWHITE) //P27087
  - F-PLACE1001257//RING CANAL PROTEIN (KELCH PROTEIN).//4.1e-24:125:46//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- 40 F-PLACE1001272//HYPOTHETICAL PROTEIN IN KSGA 3'REGION (ORF L5) (FRAGMENT).//1.0:24:45//MYC-OPLASMA CAPRICOLUM //P43040
  - F-PLACE1001279//CYTOTOXIN 3 (CYTOTOXIN V-II-3),//0.98;31;41//NAJA MOSSAMBICA (MOZAMBIQUE CO-BRA).//P01470
- F-PLACE1001280//PROCOLLAGEN ALPHA 1(II) CHAIN PRECURSOR [CONTAINS: CHONDROCALCIN].// 45 0.0051:156:32//MUS MUSCULUS (MOUSE).//P28481
  - F-PLACE1001294//GAMETOGENESIS EXPRESSED PROTEIN GEG-154.//3.7e-56:109:93//MUS MUSCULUS (MOUSE) //P50636
    - F-PLACE1001304//ZINC FINGER PROTEIN 35 (ZFP-35),//3.2e-30;75:57//MUS MUSCULUS (MOUSE),//P15620 F-PLACE1001311//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.7e-31:66:66//HOMO SAPIENS (HUMAN).// P39189
  - F-PLACE1001323

- F-PLACE1001351//REV PROTEIN (ANTI-REPRESSION TRANSACTIVATOR PROTEIN) (ART/TRS).//0.11:66: 27//SIMIAN IMMUNODEFICIENCY VIRUS (AGM155 ISOLATE) (SIV-AGM).//P27971
- F-PLACE1001366//SHORT NEUROTOXIN 2 (TOXIN CM-14) (TOXIN V-N-12),//0.070;18;33//NAJA HAJE ANNU-55 LIFERA (BANDED EGYPTIAN COBRA), //P01422
  - F-PLACE1001377//DISINTEGRIN TRIGRAMIN BETA (PLATELET AGGREGATION ACTIVATION INHIBITOR).// 4.9e-06:50:46//TRIMERESURUS GRAMINEUS (INDIAN GREEN TREE VIPER) (GREEN HABU SNAKE).// P17495

- F-PLACE1001383//M PROTEIN SEROTYPE 49 PRECURSOR://0.080:136:24//STREPTOCOCCUS PYO-GENES //P16947
- F-PLACE1001384
- F-PLACE1001387//EPIDERMAL GROWTH FACTOR RECEPTOR KINASE SUBSTRATE EPS8.//1.9e-22:142: 39//HOMO SAPIENS (HUMAN).//Q12929
  - F-PLACE1001395//HYPOTHETICAL 8.5 KD PROTEIN IN ASIA-MOTA INTERGENIC REGION.//0.98:67:34// BACTERIOPHAGE T4.//P22917
    - F-PLACE1001399//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.1e-32:47:74//HOMO SAPIENS (HUMAN).// P39194
- 10 F-PLACE1001412//GLYCOPHORIN C (PAS-2') (GLYCOPROTEIN BETA) (GLPC) (GLYCOCONNECTIN) (SIALOGLYCOPROTEIN D) (GLYCOPHORIN D) (GPD).//0.00021:125:36//HOMO SAPIENS (HUMAN).//P04921 F-PLACE1001414//CHYMOTRYPSIN/ELASTASE ISOINHIBITORS 2 TO 5.//0.99:37:35//ASCARIS SUUM (PIG ROUNDWORM) (ASCARIS LUMBRICOIDES).//P07852
  - F-PLACE1001440//PROLINE-RICH PEPTIDE P-B.//0.35:16:50//HOMO SAPIENS (HUMAN).//P02814
- 15 F-PLACE1001456//RELAXIN.//0.48:38:36//BALAENOPTERA ACUTOROSTRATA (MINKE WHALE) (LESSER RORQUAL).//P11184
  - F-PLACE1001468//HYPOTHETICAL PROTEIN MJ0602.//0.10:86:32//METHANOCOCCUS JANNASCHII.// O58019
  - F-PLACE1001484//HYPOTHETICAL 7.5 KD PROTEIN IN DNAC-RPLI INTERGENIC REGION.//1.0:47:34//BA-CILLUS SUBTILIS //P37480
    - F-PLACE1001502//COLLAGEN 1(X) CHAIN PRECURSOR.//0.00029:118:34//BOS TAURUS (BOVINE).//P23206 F-PLACE1001503//HYPOTHETICAL 77.3 KD PROTEIN T05G5.8 IN CHROMOSOME III.//2.2e-07:107:30// CAENORHABDITIS ELEGANS.//P34561
- F-PLACE1001517//SMALL PROTEIN INHIBITOR OF INSECT ALPHA-AMYLASES 2 (SI ALPHA-2).//0.56:22:45// 25 SORGHUM BICOLOR MILO (SORGHUM), //P21924
  - F-PLACE1001534//PUTATIVE GENE PROTEIN 54.//0.43:44:40//BACTERIOPHAGE SP01.//O48408
    - F-PLACE1001545//HYPOTHETICAL 7.9 KD PROTEIN IN CELF-KATE INTERGENIC REGION.//0.99:70:32//ES-CHERICHIA COLL/P37795
- F-PLACE1001551//CHLOROPLAST 50S RIBOSOMAL PROTEIN L32.//1.0:66:28//MARCHANTIA POLYMOR-PHA (LIVERWORT),//P12196
- F-PLACE1001570//SYNAPTONEMAL COMPLEX PROTEIN 1 (SCP-1 PROTEIN).//0.024:120:27//HOMO SAPI-
  - ENS (HUMAN).//Q15431 F-PLACE1001602//CCR4-ASSOCIATED FACTOR 1 (CAF1).//1.1e-30:90:78//MUS MUSCULUS (MOUSE).//
- Q60809 35 F-PLACE1001603//ACIDIC PROLINE-RICH PROTEIN PRECURSOR (CLONE PRP25) (FRAGMENT),//0.054: 77:33//RATTUS NORVEGICUS (RAT).//P10164
  - F-PLACE1001608

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- F-PLACE1001610//PROBABLE E4 PROTEIN.//0.90:58:29//HUMAN PAPILLOMAVIRUS TYPE 28.//P51896 F-PLACE1001611//METALLOTHIONEIN-IG (MT-1G).//0.35:30:40//HOMO SAPIENS (HUMAN).//P13640
- F-PLACE1001632//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2).//3.6e-28:144:43//HOMO SA-40 PIENS (HUMAN).//P51523
  - F-PLACE1001634//PHOTOSYSTEM II REACTION CENTRE N PROTEIN. // 1.0:36:41//CYANIDIUM CALDARIUM (GALDIERIA SULPHURARIA).//O19926
- F-PLACE1001640//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN),//0.24:47:38//HUMAN IMMU-45 NODEFICIENCY VIRUS TYPE 1 (NDK ISOLATE) (HIV-1).//P18804
  - F-PLACE1001672/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.0:27:66//HOMO SAPIENS (HUMAN).// P39188
    - F-PLACE1001691//HYPOTHETICAL 15.5 KD PROTEIN IN PIK1-POL2 INTERGENIC REGION.//0.40:81:33// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST). //P53842
- F-PLACE1001692//S-ACYL FATTY ACID SYNTHASE THIOESTERASE, MEDIUM CHAIN (EC 3.1.2.14) 50 (THIOESTERASE II).//8.3e-41:103:55//RATTUS NORVEGICUS (RAT).//P08635
  - F-PLACE1001705 F-PLACE1001716//HYPOTHETICAL 138.5 KD PROTEIN C17H9.01 IN CHROMOSOME L//6.1e-07:157:29// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST), //O13798

F-PLACE1001720

- F-PLACE1001729//PROLINE-RICH PROTEIN MP-3 (FRAGMENT),//6.5e-05;196:32//MUS MUSCULUS (MOUSE).//P05143
- F-PLACE1001739//NEUROFILAMENT TRIPLET M PROTEIN (160 KD NEUROFILAMENT PROTEIN) (NF-M).//

0.00050:213:23//RATTUS NORVEGICUS (RAT).//P12839

F-PLACE1001740//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.4e-17:90:56//HOMO SAPIENS (HUMAN).//

F-PLACE1001745//HYPOTHETICAL PROTEIN KIAA0125.//0.96:38:36//HOMO SAPIENS (HUMAN).//Q14138 F-PLACE1001746//CONGLUTIN DELTA-2 SMALL CHAIN.//0.98:23:43//LUPINUS ANGUSTIFOLIUS (NARROW-

LEAVED BLUE LUPINE).//P09930
F-PLACE1001748//HYPOTHETICAL 99.0 KD PROTEIN SPBC119.17.//2.9e-28:167:38//SCHIZOSACCHARO-MYCES POMBE (FISSION YEAST).//O42908

F-PLACE1001756/!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//9.2e-43:126:77//HOMO SAPIENS (HU-MAN).//P39189

MAINI, III 53163
F-PLACE1001761I/50S RIBOSOMAL PROTEIN L35.//0.26:42:38//HELICOBACTER PYLORI (CAMPYLO-BACTER PYLORI).//P56057

F-PLACE1001771//TRANSIENT-RECEPTOR-POTENTIAL LIKE PROTEIN.//4.8e-35:223:40//DROSOPHILA MELANOGASTER (FRUIT FLY).//P48994

F-PLACE1001781//HYPOTHETICAL 71.1 KD PROTEIN IN DSK2-CAT8 INTERGENIC REGION.//9.5e-41:194: 46//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q03262

F-PLACE1001799

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F-PLACE1001810

F-PLACE1091817//SUCCINYL-COA LIGASE [GDP-FORMING], BETA-CHAIN PRECURSOR (EC 6.2.1.4) (SUCCINYL-COA SYNTHETASE, BETA CHAIN) (SCS-BETA).//2.8e-40:115:61//NEOCALLIMASTIX FRONTALIS (RUMEN FUNGUS)./PS3587

F-PLACE1001821

F-PLACE1001844//IG KAPPA CHAIN V-I REGION (HAU).//0.59:89:35//HOMO SAPIENS (HUMAN).//P01600

25 F-PLACE1001869//MPA43 PROTEIN.//3.5e-14:153:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P53583

F-PLACE1001897/ILIGATOXIN A./1.0-43:27//PHORADENDRON LIGA (ARGENTINE MISTLETOE).//P01540 F-PLACE1001912//LONG NEUROTOXIN 2 (TOXIN C).//0.57:44:45//ASTROTIA STOKESI (STOKES'S SEA SNAKE) (DISTEIRA STOKESI).//P01391

F-PLACE1001920//LATE GENES ACTIVATOR (EARLY PROTEIN GP4) (GPF).//0.89:75:29/BACTERIOPHAGE NF.//P09877

F-PLACE1001928

F-PLACE1001983//IMMEDIATE-EARLY PROTEIN IE180.//0.0049:51:45//PSEUDORABIES VIRUS (STRAIN KA-PLAN) (PRV).//P33479

35 F-PLACE1001989//PUTATIVE AMIDASE (EC 3.5.1.4).//8.9e-08:125:36//MORAXELLA CATARRHALIS.//Q49091 F-PLACE1002004

F-PLACE1002046//LIGATIN (FRAGMENT).//1.6e-84:191:84//MUS MUSCULUS (MOUSE).//Q61211 F-PLACE1002052

F-PI ACE 1002066

40 F-PLACE1002072//ANTER-SPECIFIC PROLINE-RICH PROTEIN APG PRECURSOR.//0.16:77:31//ARABIDOP-SIS THALIANA (MOUSE-EAR CRESS).//P40602

F-PLACE1002073/HYPOTHETICAL 118.2 KD PROTEIN F43C1.1 IN CHROMOSOME III.//4.0e-11:174:28//CAENORHABDITIS ELEGANS.//Q09564

F-PLACE1002090//SIGNAL RECOGNITION PARTICLE 72 KD PROTEIN (SRP72).//2.8e-57:112:99//HOMO SA-PIENS (HUMAN).//076094

F-PLACE1002115/P8 MTCP-1 PROTEIN (MATURE T-CELL PROLIFERATION-1 TYPE A) (MTCP-1 TYPE A) (PBMTCP1)//1.0:49:30/MUS MUSCULUS (MOUSE)//Q61908
F-PLACE1002119//T-LYMPHOCYTE ACTIVATED PROTEIN (CYCLOHEXIMIDE-INDUCED) (CHX1) (IMMEDI-

ATE EARLY RESPONSE 2 PROTEIN).//2.7e-11:118:36//MUS MUSCULUS (MOUSE).//P17950

F-PLACE1002140/HYPOTHETICAL 12.3 KD PROTEIN IN MOBL 3'REGION (ORF 4).//0.0086:39.46//THIOBA-CILLUS FERROOXIDANS.//P20088 E-PLACE1002150

F-PLACE1002157//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!/2.4e-34:56:82//HOMO SAPIENS (HUMAN).// P39189

55 F-PLACE1002163//NEUROTOXIN 1.//1.0:17:52//CENTRUROIDES SCULPTURATUS (BARK SCORPION).// P01492

F-PLACE1002170

F-PLACE1002171//TRANSCRIPTION REGULATORY PROTEIN SWI3 (SWI/SNF COMPLEX COMPONENT

- SWI3) (TRANSCRIPTION FACTOR
- TYE2).//0.00023:179:23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32591
- F-PLACE1002205//HYPOTHETICAL 13.5 KD PROTEIN IN MOB1-SGA1 INTERGENIC REGION.//0.77:21:47// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40490
- F-PLACE1002213//HISTONE H4 (FRAGMENT).//0.62;31:32//BLEPHARISMA JAPONICUM.//P80738
  - F-PLACE1002227//HYPOTHETICAL 7.9 KD PROTEIN IN FIXW 5'REGION.//0.41:49:36//RHIZOBIUM LEGUMI-NOSARUM.//P14310
    - F-PLACE1002256//CYTOCHROME B (EC 1.10.2.2).//0.61:95:29//CAENORHABDITIS ELEGANS.//P24890
- F-PLACE1002259//HYPOTHETICAL 9.2 KD PROTEIN IN SPS1-QCR7 INTERGENIC REGION.//0.99:22:45//
  SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P56508
  - F-PLACE1002319//HYPOTHETICAL 56.6 KD PROTEIN IN URE2-SSU72 INTERGENIC REGION.//0.91:18:72// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53867
  - F-PLACE1002342//HYPOTHETICAL PROTEIN C16.//1.0:53:32//SWINEPOX VIRUS (STRAIN KASZA) (SPV).// P32219
- 15 F-PLACE1002395//CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS).//6.4e-05:127:37//PLASMODIUM VIVAX.//P08677
  - F-PLACE1002399
    - F-PLACE1002433//DYNACTIN, 150 KD ISOFORM (150 KD DYNEIN-ASSOCIATED POLYPEPTIDE) (DP-150) (DAP-150) (P150-GLUED) //0.00094:182:25//RATTUS NORVEGICUS (RAT) //P28023
- F-PLACE1002437//ATP-BINDING CASSETTE TRANSPORTER 1.//4.5e-19:62:77//MUS MUSCULUS (MOUSE).//P41233
  - (MOUSE).//P41233
    F-PLACE1002438//HYPOTHETICAL 141.5 KD ZINC FINGER PROTEIN IN TUB1-CPR3 INTERGENIC RE-
    - GION.//0.014:63:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04545 F-PLACE1002450//OOCYTE ZINC FINGER PROTEIN XLCOF6 (FRAGMENT).//3.9e-28:159:38//XENOPUS
- 25 LAEVIS (AFRICAN CLAWED FROG).//P18749 F-PLACE1002465/LARIAT DEBRANCHING ENZYME (EC 3.1.--).//0.0014:148:28//SCHIZOSACCHAROMY-CES POMBE (FISSION YEAST).//013765
- F-PLACE1002474//FIBRILLIN 2 PRECURSOR.//2.1e-24:203:33//MUS MUSCULUS (MOUSE).//Q61555
- F-PLACE1002477//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//0.15:65:41//HOMO SAPIENS (HUMAN).// 90 P39193
- F-PLACE1002493//SEMENOGELIN II PRECURSOR (SGII).//1.0:72:31//MACACA MULATTA (RHESUS MACAQUE).//Q95196
  - F-PLACE1002499//HYPOTHETICAL 39.3 KD PROTEIN C02B8.6 IN CHROMOSOME X.//2.9e-11:67:35// CAENORHABDITIS ELEGANS.//Q11096
- 35 F-PLACE1002500//COBALT-ZINC-CADMIUM RESISTANCE PROTEIN CZCD (CATION EFFLUX SYSTEM PRO-TEIN CZCD).//8.4e-11:143:32//ALCALIGENES EUTROPHUS.//P13512
  - F-PLACE1002514//HYPOTHETICAL 8.1 KD PROTEIN IN SPEA-METK INTERGENIC REGION (071).//1.0:15: 60//ESCHERICHIA COLI.//P46878
  - F-PLACE1002529
- 40 F-PLACE1002532//HOMEOBOX PROTEIN DLX-5.//1.1e-76:183:81//MUS MUSCULUS (MOUSE).//P70396 F-PLACE1002537//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.6e-18:51:86//HOMO SAPIENS (HUMAN).//
  - F-PLACE1002571//ACTIN-LIKE PROTEIN 13E.//6.0e-56:140:47//DROSOPHILA MELANOGASTER (FRUIT FLY) //P45890
- 45 F-PLACE1002578
  - F-PLACE1002583
  - F-PLACE1002591//CORONIN-LIKE PROTEIN P57.//5.5e-26:78:69//BOS TAURUS (BOVINE).//Q92176
  - F-PLACE1002598
  - F-PLACE1002604
- 50 F-PLACE1002625//HYPOTHETICAL 180.2 KD PROTEIN IN FAA4-HOR7 INTERGENIC REGION.//6.4e-08:193: 23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04781
  - F-PLACE1002655//ADSEVERIN (GELSOLIN-LIKE PROTEIN).//7.1e-100:210:89//MUS MUSCULUS (MOUSE).//
- F-PLACE1002665/MOBILIZATION PROTEIN MOBS.//0.35.60.30/THIOBACILLUS FERROOXIDANS.//P20086
  55 F-PLACE1002685/ACTIN BINDING PROTEIN.//0.052:115:29//SACCHAROMYCES EXIGUUS (YEAST).//
  - F-PLACE1002714//CIS-GOLGI MATRIX PROTEIN GM130.//1.8e-06:214:30//RATTUS NORVEGICUS (RAT).// Q62839

- F-PLACE1002722//THROMBIN RECEPTOR PRECURSOR.//2.0e-19:134:38//XENOPUS LAEVIS (AFRICAN CLAWED FROG) //P47749
- F-PLACE1002768//FOLLICLE STIMULATING HORMONE RECEPTOR PRECURSOR (FSH-R) (FOLLITROPIN RECEPTOR) (FRAGMENT).//0.43:40:35//MUS MUSCULUS (MOUSE).//P35378
- F-PLACE1002772

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- F-PLACE1002775//CENTROMERE/MICROTUBULE BINDING PROTEIN CBF5 (CENTROMERE-BINDING FAC-TOR 5) (NUCLEOLAR PROTEIN CBF5).//4.8e-07:96:29//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O14007
- F-PLACE1002782//COBALT-ZINC-CADMIUM RESISTANCE PROTEIN CZCD (CATION EFFLUX SYSTEM PRO-TEIN CZCD\//1.1e-07:114:35//ALCALIGENES EUTROPHUS //P13512
- F-PLACE1002794//CUTICLE COLLAGEN 12 PRECURSOR.//0.0068:98:39//CAENORHABDITIS ELEGANS.// P20630
- F-PLACE1002811//CYCLIN-DEPENDENT KINASE 6 INHIBITOR (P18-INK6) (CYCLIN-DEPENDENT KINASE 4 INHIBITOR C) (P18-INK4C).//1.1e-09:137:34//MUS MUSCULUS (MOUSE).//Q60772
- 15 F-PLACE1002815//C-HORDEIN (CLONE PC HOR1-3) (FRAGMENT).//0.46:35:42//HORDEUM VULGARE (BAR-LEY).//P17991
  - F-PLACE1002816//HYPOTHETICAL PROTEIN KIAA0288 (HA6116).//1.0e-86:201:74//HOMO SAPIENS (HU-MAN).//P56524
  - F-PLACE1002834//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//1.6e-30:54:96//HOMO SAPI-ENS (HUMAN).//P51522
  - F-PLACE1002839//METALLOTHIONEIN-I (MT-I).//1.0:43:37//MUS MUSCULUS (MOUSE).//P02802
    - F-PLACE1002851//BOWMAN-BIRK TYPE PROTEINASE INHIBITOR (VAI).//0.77:35:37//VICIA ANGUSTIFOLIA (COMMON VETCH).//P01065
- F-PLACE1002853//HYPOTHETICAL 7.9 KD PROTEIN IN PE 5'REGION (ORF1).//1.0:18:55//LYMANTRIA DIS-PAR MULTICAPSID NUCLEAR POLYHEDROSIS VIRUS (LDMNPV).//P36866
  - F-PLACE1002881//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.1e-27:91:70//HOMO SAPIENS (HUMAN).//
    - F-PLACE1002908//HYPOTHETICAL 33.8 KD PROTEIN R10E11.4 IN CHROMOSOME III.//2.0e-31:148:46// CAENORHABDITIS ELEGANS.//P34548
- 30 F-PLACE1002941//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.6e-11:40:85//HOMO SAPIENS (HUMAN).// P39195
  - F-PLACE1002962//ENDOTHELIN-1 PRECURSOR (ET-1) (FRAGMENT).//0.90:38:36//CANIS FAMILIARIS (DOG).//P13206
    - F-PLACE1002968//TOXIN IV-5 PRECURSOR (TITYUSTOXIN) (FRAGMENT).//0.97:26:38//TITYUS SERRULA-TUS (BRAZILIAN SCORPION).//P01496
    - F-PLACE1002991//PUTATIVE AMIDASE (EC 3.5.1.4).//3.3e-20:120:41//METHANOCOCCUS JANNASCHII.// Q58560
    - F-PLACE1002993//HYPOTHETICAL 17.8 KD PROTEIN IN SMPA-SMPB INTERGENIC REGION (F158).// 0.00045:93:23//ESCHERICHIA COLI.//P52121
- 40 F-PLACE1002996//PUTATIVE REGULATORY PROTEINTSC-22 (TGFB STIMULATED CLONE 22 HOMOLOG).// 0.17:91:29//GALLUS GALLUS (CHICKEN).//Q91012
  - F-PLACE1003025/ISUPPRESSOR PROTEIN SRP40.//0.0079:214:24//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//P32563 F-PLACE1003027//HYPOTHETICAL 128.6 KD PROTEIN ZK1098.10 IN CHROMOSOME III.//1.3e-49:167:63//
  - CAENORHABDITIS ELEGANS.//P34609
    - F-PLACE1003044//SPORE COAT PROTEIN D.//0.97:24:45//BACILLUS SUBTILIS.//P07791
    - F-PLACE1003045
      - E DI ACE1000040
    - F-PLACE1003100//HEP27 PROTEIN (PROTEIN D).//3.9e-51:188:57//HOMO SAPIENS (HUMAN).//Q13268 F-PLACE1003108
- 50 F-PLACE1003108 F-PLACE1003136
  - F-PLACE1003145//BUTYROPHILIN PRECURSOR (BT).//0.00024:170:24//BOS TAURUS (BOVINE).//P18892
- F-PLACE1003153//HUNCHBACK PROTEIN (FRAGMENT).//1.0:32:37//LOCUSTA MIGRATORIA (MIGRATORY LOCUST).//Q01777
- 55 F-PLACE1003174/JUBIQUITIN-CONJUGATING ENZYME E2-18 KD (EC 6.3.2.19) (UBIQUITIN- PROTEIN LIGASE) (UBIQUITIN CARRIER PROTEIN) (PM42).//6.3e-05.54:38//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P42743
  - F-PLACE1003176//HYPOTHETICAL 62.3 KD PROTEIN IN PCS60-ABD1 INTERGENIC REGION.//0.24;74;36//

SACCHAROMYCES CEREVISIAE (BAKER'S YEAST), I/P38319

F-PLACE1003190//SOF1 PROTEIN.//1.0e-52:158:41//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P33750

#### F-PLACE1003200

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F-PLACE1003205//SPERM PROTAMINE P1.//0.074;20;45//CAENOLESTES FULIGINOSUS.//P42131

F-PLACE1003238//PROBABLE G PROTEIN-COUPLED RECEPTOR KIAA0001.//0.013:20:55//HOMO SAPIENS (HUMAN).//015391

F-PLACE1003249/HYPOTHETICAL PROTEIN KIAA0125.//0.98-48:37//HOMO SAPIENS (HUMAN).//Q14138 F-PLACE1003256//OMEGA-CONOTOXINS GVII, GVIB AND GVIC PRECURSOR (SHAKER PEPTIDE).//0.84: 53:39//CONUS GEOGRAPHUS (GEOGRAPHY CONE).//P01522

F-PLACE1003258//EARLY EMBRYOGENESIS ZYG-11 PROTEIN.//4,1e-18:70:47//CAENORHABDITIS ELE-GANS.//P21541

F-PLACE1003298//SPECTRIN BETA CHAIN, ERYTHROCYTE.//0.063:160:24//HOMO SAPIENS (HUMAN).// P11277

F-PLACE1003302//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1),//9.4e-69:84:94//HOMO SAPI-ENS (HUMAN),//P51522

F-PLACE1003334//NUCLEOBINDIN PRECURSOR (NUCB1) (BONE 63 KD CALCIUM-BINDING PROTEIN).// 0.029:125:24//RATTUS NORVEGICUS (RAT).//Q63083

F-PLACE1003342//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.97:44:40//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01643

F-PLACE1003343//GENE 11 PROTEIN.//1.0:37:37//SPIROPLASMA VIRUS SPV1-R8A2 B.//P15902

 $\label{eq:fplace} F-PLACE1003353//SH2/SH3 ADAPTOR CRK (ADAPTER MOLECULE CRK) (CRK2).//6.4e-05:69:40//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P87378$ 

F-PLACE1003361//!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.6e-23:66:75//HOMO SAPIENS (HUMAN).// P39192

F-PLACE1003368/ISMALL PROLINE-RICH PROTEIN 2-1/I/0.62:19:57/I/HOMO SAPIENS (HUMAN) /I/P35326 F-PLACE1003369/IA-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//4.3e-06:102:42//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST) /I/P32323

F-PLACE1003373//PROTEIN Q300.//0.042:29:37//MUS MUSCULUS (MOUSE).//Q02722

30 F-PLACE1003375/OLFACTORY RECEPTOR 11 (M49) (FRAGMENT).//0.99:46:34//MUS MUSCULUS (MOUSE).//060890 F-PLACE1003383

F-PLACE1003394//RAS-RELATED PROTEIN RAB-14.//2.8e-80:166:89//RATTUS NORVEGICUS (RAT).//

### F-PLACE1003401

F-PLACE1003420//PUTATIVE MITOCHONDRIAL CARRIER YIL006W.//8.1e-17:138:37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40556

# F-PLACE1003454

F-PLACE1003478

40 F-PLACE1003493//ENDOTHELIAL CELL MULTIMERIN PRECURSOR.//3.4e-11:123:32//HOMO SAPIENS (HU-MAN).//Q13201

F-PLACE1003516//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.0e-32:68:76//HOMO SAPIENS (HU-MAN).//P08547

F-PLACE1003519//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//9.2e-17:77:50//HOMO SAPIENS (HUMAN).// P39188

F-PLACE1003521//HYPOTHETICAL BAMHI-ORF9 PROTEIN.//1.0:38:42//FOWLPOX VIRUS (ISOLATE HP-438 [MUNICH]).//P14366

F-PLACE1003528//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.96:32:40//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P03931

F-PLACE1003537//CEF PROTEIN //0 92:47:29//BACTERIOPHAGE T4 //Q01436

F-PLACE1003553

F-PLACE1003566//HYPOTHETICAL BAMHI-ORF9 PROTEIN.//1.0:32:34//FOWLPOX VIRUS (ISOLATE HP-438 [MUNICH]).//P14366

E-PLACE1003575

F-PLACE1003583//PROBABLE E5 PROTEIN.//0.16:64:31//HUMAN PAPILLOMAVIRUS TYPE 35.//P27226 F-PLACE1003584

F-PLACE1003592//EXCISIONASE.//0.26:19:52//BACTERIOPHAGE PHI-80.//P05998

F-PLACE1003593//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:42:30//OVIS ARIES (SHEEP).//

### O78751

- F-PLACE1003596//OLIGOSACCHARYL TRANSFERASE STT3 SUBUNIT HOMOLOG.//6.3e-87:238:67// CAENORHABDITIS ELEGANS.//P46975
- F-PLACE1003602//HYPOTHETICAL 11.0 KD PROTEIN IN FAA3-MAS3 INTERGENIC REGION.//8.4e-17:98:42// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40554
  - F-PLACE1003605//HAP5 TRANSCRIPTIONAL ACTIVATOR.//2.0e-09:82:35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q02516
  - F-PLACE1003611//PANCREATIC SECRETORY TRYPSIN INHIBITOR://0.99:32:43//CANIS FAMILIARIS (DOG).//P04542
- 10 F-PLACE1003618/LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.7e-65:229:58//HOMO SAPIENS (HU-MAN).//P08547
  - F-PLACE1003628/J0S RIBOSOMAL PROTEIN S20 (FRAGMENT) //1.0:56:26//PROTEUS MIRABILIS.//P42275 F-PLACE1003638//PROTEIN Q300./0.079.41:39//MUS MUSCULUS (MOUSE).//Q02722 F-PLACE1003669//TRICHOHYALIN.//2.9e-07:160:30//OVIS ARIES (SHEEP).//P22733
- 15 F-PLACE1003704//SPLICING FACTOR, ARGININE/SERINE-RICH 4 (PRE-MRNA SPLICING FACTOR SRP75), //3.3e-16:98-40//HOMO SAPIENS (HUMAN), //008170
  - F-PLACE1003709//HYPOTHETICAL 59.5 KD PROTEIN IN CCT3-CCT8 INTERGENIC REGION.//2.8e-07:128: 27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47074
  - F-PLACE1003711//ALPHA/BETA-GLIADIN PRECURSOR (PROLAMIN) (CLASS A-IV).//5.0e-05:88:30//TRITI-CUM AESTIVUM (WHEAT).//P04724
    - F-PLACE1003723//TYROSINE-PROTEIN KINASE SRM (EC 2.7.1.112) (PTK70).//6.0e-06:98:36//MUS MUSCU-LUS (MOUSE).//062270
    - F-PLACE1003738//OOCYTE ZINC FINGER PROTEIN XLCOF6 (FRAGMENT).//2.5e-45:147:46//XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P18749
- F-PLACE1093760/ICYTOCHROME B (EC 1.10.2.2).//0.91:49:34/TRYPANOSOMA BRUCEI BRUCEI.//P00164 F-PLACE1003762//METALLOTHIONEIN-LIKE PROTEIN TYPE 2.//0.98:28:32/MALUS DOMESTICA (APPLE) (MALUS SYLVESTRIS).//0.24058
  - F-PLACE1003768/JLINE-1 REVERSE TRANSCRIPTASE HOMOLOG J/8.5e-19:123:37//HOMO SAPIENS (HU-MAN).//P08547
- 30 F-PLACE1003771

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- F-PLACE1003783//SRY-RELATED PROTEIN ADW2 (FRAGMENT).//1.0:29:37//ALLIGATOR MISSISSIPPIEN-SIS (AMERICAN ALLIGATOR).//P40634
  - F-PLACE1003784//HYPOTHETICAL 98.1 KD PROTEIN IN SPX19-GCR2 INTERGENIC REGION.//1.2e-13:199: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40164
- F-PLACE1003795//EC PROTEIN I/II (ZINC-METALLOTHIONEIN CLASS II).//0.67:53:30//TRITICUM AESTIVUM (WHEAT).//P30569
  - F-PLACE1003833/METHIONYL-TRNA FORMYLTRANSFERASE (EC 2.1.2.9).//0.99:158:28//THERMUS AQUATICUS (SUBSP. THERMOPHILUS).//P43523
  - F-PLACE1003850
- F-PLACE1003858/I-UNCHBACK PROTEIN (FRAGMENT)/IO 37:28:42/I/LITHOBIUS FORFICATUS /IQQ2030 F-PLACE1003864/I/QUTER MEMBRANE LIPOPROTEIN LOLB PRECURSOR./IO 0046-116:31/IACTINOBACIL-LUS ACTINOMYCETEMCOMITANS (HAEMOPHILUS ACTINOMYCETEMCOMITANS)//IO52727 F-PLACE1003870
- F-PLACE1003885//POLY(A) POLYMERASE (EC 2.7.7.19) (PAP) (POLYNUCLEOTIDE ADENYLYLTRANS-FERASE) (FRAGMENT).//1.6e-92:166:75//HOMO SAPIENS (HUMAN).//P51003
- F-PLACE1003886//MMEDIATE-EARLY PROTEIN IE180.//0.54:96:34//PSEUDORABIES VIRUS (STRAIN INDI-ANA-FUNKHAUSER / BECKER) (PRV).//P11675
  - F.PLACE1003888/I-PHOSPHATIDYLINOSITOL.4,5-BISPHOSPHATE PHOSPHODIESTERASE DELTA 1 (EC 3.1.4.11) (PLC-DELTA-1) (PHOSPHOLIPASE C-DELTA-1) (PLC-III) (FRAGMENT).///8.8e-54:260:46//BOS TAU-RUS (BOVINE).//P10895
- F-PLACE1003892//PROBABLE E5 PROTEIN //1.0-13-61/H/UMAN PAPILLOMA/RUS TYPE 18.//P06792
  F-PLACE1003980//BETA-FRUCTOFURANOSIDASE, SOLUBLE 1905EVYME I (EG 2.2.126) (SUCROSE-6-PHOSPHATE HYDROLASE) (INVERTASE) (FRAGMENTS)//0.58-49:36//DAUCUS CAROTA (CARROT)///
- 55 F-PLACE1003903//CTP SYNTHASE (EC 6.3.4.2) (UTP--AMMONIA LIGASE) (CTP SYNTHETASE).//3.8e-52:92: 85//HOMO SAPIENS (HUMAN).//P17812
  - F-PLACE1003915//PROBABLE ARGINYL-TRNA SYNTHETASE, CYTOPLASMIC (EC 6.1.1.19) (ARGININE-TRNA LIGASE) (ARGRS).//2.6e-26:202:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q05506

- F-PLACE1003923//HISTIDYL-TRNA SYNTHETASE (EC 6.1.1.21) (HISTIDINE--TRNA LIGASE) (HISRS).//0.94: 65:29//STREPTOCOCCUS FOLISIMILIS //P30053
- F-PLACE1003932//HYPOTHETICAL 17.3 KD PROTEIN IN SEC15-SAP4 INTERGENIC REGION.//0.098:79:31// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53074
- F-PLACE1003936

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- F-PLACE1003968//5'-AMP-ACTIVATED PROTEIN KINASE, GAMMA-1 SUBUNIT (AMPK GAMMA-1 CHAIN).//
  4 7e-68:164:78//RATTUS NORVEGICUS (RAT).//P80385
- F-PLACE1004103//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.9e-14:60:73//HOMO SAPIENS (HUMAN).//
- 10 F-PLACE1004104//EXOCYST COMPLEX COMPONENT SEC5.//0.020:202:20//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P89102
  - F-PLACE1004114//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.1e-15:69:60//HOMO SAPIENS (HUMAN).// P39188
  - F-PLACE1004118//REGULATORY PROTEIN E2.//0.73:58:36//CANINE ORAL PAPILLOMAVIRUS (COPV).//
  - F-PLACE1004128//GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT 4 (TRANSDUCIN BETA CHAIN 4),//7.7e-62:108:100//MUS MUSCULUS (MOUSE).//P29387
    - F-PLACE1004149//PROBABLE NUCLEAR ANTIGEN.//0.0011:73:42//PSEUDORABIES VIRUS (STRAIN KAP-LAN) (PRV).//P33485
- F-PLACE1004156//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//0.00061:39:48//OWENIA FUSI-FORMIS.//P21260
  - F-PLACE1004161//PLASMINOGEN-BINDING PROTEIN PAM PRECURSOR (FRAGMENT).//0.033:108:27// STREPTOCOCCUS PYOGENES //P44054
  - F-PLACE1004183//HYPOTHETICAL 64.3 KD PROTEIN IN CDC12-ERP5 INTERGENIC REGION.//4.0e-07:146: 35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38817
  - F-PLACE1004197//BUTYROPHILIN PRECURSOR (BT).//5.9e-11:208:27//MUS MUSCULUS (MOUSE).//
    - F-PLACE1004203//PROTEIN A39.//8.5e-18:139:33//VACCINIA VIRUS (STRAIN COPENHAGEN).//P21062
      F-PLACE1004242//PHOTOSYSTEM II REACTION CENTRE J PROTEIN.//1.0:28:42//PISUM SATIVUM (GAR-
- 30 DEN PEA).//P13555
  F-PLACE1004256//MYOSIN HEAVY CHAIN D (MHC D).//0.73:134:25//CAENORHABDITIS ELEGANS.//P02567
  F-PLACE1004256//MYOSIN HEAVY CHAIN D (MHC D).//0.73:134:25//CAENORHABDITIS ELEGANS.//P02567
  F-PLACE1004257/MYOSIN HEAVY CHAIN D (MHC D).//0.73:134:25//CAENORHABDITIS ELEGANS.//P02567
  F-PLACE1004257/MYOSIN HEAVY CHAIN D (MHC D).//0.73:134:25//CAENORHABDITIS ELEGANS.//P02567
  F-PLACE1004257/MYOSIN HEAVY CHAIN D (MHC D).//0.73:134:25//CAENORHABDITIS ELEGANS.//P02567
  - F-PLACE:1004257/I/YPOTHETICAL PROTEIN HI0490.//0.13:75:29/I/HAEMOPHILUS INFLUENZAE./I/P44006 F-PLACE:1004258/I/COLLAGEN ALPHA 2(VIII) CHAIN (ENDOTHELIAL COLLAGEN) (FRAGMENT).//0.027:128: 35/I/HOMO SAPIENS (HUMAN)./I/25067
- 35 F-PLACE1004270//LARGE TEGUMENT PROTEIN.//1.8e-10:100:44//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03186
  - F-PLACE1004274/IH/POTHETICAL PROTEIN E-95.//0.44:61:42/IHUMAN ADENOVIRUS TYPE 2.//P/03286 F-PLACE1004277//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//0.0013:55:38//BOS TAURUS (BOVINE).// P25508
- 40 F-PLACE1004284//7 KD PROTEIN (ORF 4).//1.0:63:23//CHRYSANTHEMUM VIRUS B (CVB).//P37990
  - F-PLACE1004289/SPERM PROTAMINE P3 //0.00057:22:77/JMUS MUSCULUS (MOUSE) //062100 F-PLACE1004302I/SERINE/THREONINE PROTEIN KINASE AFSK (EC 2.7.1.-).//0.0065:148:29//STREPTOMY-CES COELICOLOR //P54741
- F-PLACE1004316//AUTOPHAGY PROTEIN APG5.//8.8e-06:117:29//SACCHAROMYCES CEREVISIAE (BAK-45 ER'S YEAST).//Q12380
  - F-PLACE1004336//COLLAGEN ALPHA 4(IV) CHAIN PRECURSOR.//0.0027:83:36//HOMO SAPIENS (HU-MAN).//P53420
    - F-PLACE1004358//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//2.9e-05:200:33//GALLUS GALLUS (CHICKEN).//P02457
- F-PLACE1004376//AXONEME-ASSOCIATED PROTEIN MST101(2).//2.4e-05:179:29//DROSOPHILA HYDEI (FRUIT FLY).//Q08696
  - F-PLACE1004384//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.6e-28:46:76//HOMO SAPIENS (HUMAN).// P39194
- F-PLACE1004388//HYPOTHETICAL 75.2 KD PROTEIN IN ACS1-GCV3 INTERGENIC REGION.//5.7e-34:202: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39722
  - F-PLACE1004405/INEURAMINYLLACTOSE-BINDING HEMAGGLUTININ (N-ACETYLNEURAMINYLLAC-TOSE- BINDING FIBRILLAR HEMAGGLUTININ RECEPTOR-BINDING SUBUNIT) (NLBH) (FLAGELLAR SHEATH AOHESIN) (ADHESIN A) (FRAGMENT)://i0.937:433/i/HELICOBACTER ACINONYX/i/047947

- F-PLACE1004425//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.81:70:42//HOMO SAPIENS (HUMAN).// P39195
- F-PLACE1004428//PRISTANOYL-COA OXIDASE (EC 1.3.3.-).//1.9e-31:203:39//RATTUS NORVEGICUS (RAT).//O63448
- F-PLÁCE1004437//SOCITRATE DEHYDROGENASE (NAD), MITOCHONDRIAL SUBUNIT BETA PRECURSOR (EC 1.1.1.41) (ISOCITRIC DEHYDROGENASE) (NAD+S-PECIFIC ICDH) (FRAGMENT)/4.2e-93:140:1001/MACACA FASCICULARIS (CRAB EATING MACAQUE) (CYYDMOLGUS MONKEY)/028479
  - F-PLACE1004451//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00013:40:62//HOMO SAPIENS (HUMAN).//
- F-PLACE1004460//MATERNAL TUDOR PROTEIN.//0.0066:218:23//DROSOPHILA MELANOGASTER (FRUIT FLY) //P25823
  - F-PLACE1004467//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//7.8e-10:33:87//HOMO SAPIENS (HUMAN).// P39193
- F-PLACE1004471//ZINC FINGER PROTEIN 83 (ZINC FINGER PROTEIN HPF1).//7.0e-56:92:58//HOMO SAPI-15 ENS (HUMAN).//P51522
- F-PLACE1004473//HYPOTHETICAL 54.3 KD PROTEIN C23D3.03C IN CHROMOSOME I.//0.019:136:27// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09844
  - F-PLACE1004491//LYSIS PROTEIN.//0.95:53:30//BACTERIOPHAGE FR.//P19903

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- F-PLACE1004506//AUTOIMMUNOGENIC CANCER/TESTIS ANTIGEN NY-ESO-1 (LAGE-1).//0.58:66:34//HO-MO SAPIENS (HUMAN).//P78358
- F-PLACE1004510//TRANSCRIPTION INITIATION FACTOR TFIID 150 KD SUBUNIT (TAFII-150) (TAFII150).// 3.0e-07:63:46//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q24325
  - F-PLACE1004516//HYPOTHETICAL PROTEIN 5' TO ASP-RICH AND HIS-RICH PROTEINS (FRAGMENT).//
    0.95:62:29//PLASMODIUM FALCIPARUM (ISOLATE FCM17 / SENEGAL).//P14587
- 25 F-PLACE1004518//METALLOTHIONEIN 10-III (MT-10-III).//0.91:28:42//MYTILUS EDULIS (BLUE MUSSEL).// P80248
  - F-PLACE1004548//DIHYDROPYRIDINE-SENSITIVE L-TYPE, SKELETAL MUSCLE CALCIUM CHANNEL GAM-MA SUBUNIT.//0.94:75:32//ORYCTOLAGUS CUNICULUS (RABBIT).//P19518
- F-PLACE1004550//CUTICLE COLLAGEN 2.//0.90:155:31//CAENORHABDITIS ELEGANS.//P17656
  F-PLACE1004564//CLEAVAGE AND POLYADENYLATION SPECIFICITY FACTOR. 100 KD SUBUNIT (CPSF 100)
- KD SUBUNIT.//3.2e-70:121:100/IBOS TAURUS (BOVINE).//Q10568 F-PLACE1004629//PROTEIN OS-9 PRECURSOR./1.7e-10:132:36/IHOMO SAPIENS (HUMAN).//Q13438 F-PLACE1004645//TRANSCRIPTION INITIATION FACTOR IB HOMOLOG (TFIIB).//0.00036:100:30//PYRO-
- COCCUS FURIOSUS.//051731

  F-PLACE1004648//PROBABLE UDP-GALACTOPYRANOSE MUTASE (EC 5.4.99.9).//0.91:58:29//KLEBSIELLA PNEUMONIAE\_I/049481
  - F-PLACE1004658//GLUTAMATE [NMDA] RECEPTOR SUBUNIT EPSILON 4 PRECURSOR (N-METHYL D-AS-PARTATE RECEPTOR SUBTYPE 2D) (NR2D) (NMDAR2D).//0.031:134:32//MUS MUSCULUS (MOUSE).// 003391
- 40 F-PLACE1004664/HYPOTHETICAL 180.2 KD PROTEIN IN FAA4-HOR7 INTERGENIC REGION.//0.025:125: 20//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q04781
  - F-PLACE1004672//HYPOTHETICAL 36.7 KD PROTEIN C2F7:14C IN CHROMOSOME I.//7.6e-52:158:56// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST),//Q09704
- F-PLACE1004674//PROBABLE CALCIUM-BINDING PROTEIN ALG-2 (PMP41) (ALG-257),//1.4e-88:144:93//
  45 MUS MUSCULUS (MOUSE),//P12815
  - F-PLACE1004681//CCR4-ASSOCIATED FACTOR 1 (CAF1).//1.0e-34:70:100//MUS MUSCULUS (MOUSE).// Q60809
  - F-PLACE1004686//!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//3.4e-08:48:62//HOMO SAPIENS (HUMAN).// P39192
  - F-PLACE1004691//METALLOTHIONEIN (MT).//0.064:24:45//ARIANTA ARBUSTORUM.//P55946 F-PLACE1004693
  - F-PLACE1004693

    F-PLACE1004716/ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:27:37//PAN PANISCUS (PYGMY CHIMPANZEE) (BONOBO).//Q35587
- F-PLACE1004722/HYPOTHETICAL 61.5 KD PROTEIN IN CLA4-MID1 INTERGENIC REGION.//0.95:53:33//
  SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P48565
- F-PLACE1004736//NEURONAL AXONAL MEMBRANE PROTEIN NAP-22.//0.014:163:30//RATTUS NORVEGI-CUS (RAT).//005175
  - F-PLACE1004740//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/1.9e-09:37:70//HOMO SAPIENS (HUMAN).//

### P39194

F-PLACE1004743//HYPOTHETICAL 12.6 KD PROTEIN IN ALGR3 3'REGION.//0.99:72:33//PSEUDOMONAS

F-PLACE1004751//CMP-N-ACETYLNEURAMINATE-BETA-GALACTOSAMIDE-ALPHA-2.3-SIALYLTRANS-

FERASE (EC 2.4.99.-) (BETA-GALACTOSIDE ALPHA-2,3-SIALYLTRANSFERASE) (ST3GALIII) (ALPHA 2.3-ST) (GAL-NAC6S) (STZ) (SIAT4-C) (SAT-3) (ST-4).//2.2e-08:90:38//HOMO SAPIENS (HUMAN).//Q11206

F-PLACE1004773//ANKYRIN R (ANKYRINS 2.1 AND 2.2) (ERYTHROCYTE ANKYRIN).//3.2e-25:233:32//HOMO SAPIENS (HUMAN).//P16157

F-PLACE1004777//N-CHIMAERIN (NC) (N-CHIMERIN) (ALPHA CHIMERIN) (A-CHIMAERIN).//8.1e-26:210:30// RATTUS NORVEGICUS (RAT).//P30337

F-PLACE1004793//ENV POLYPROTEIN [CONTAINS: COAT PROTEIN GP52; COAT PROTEIN GP36].//0.00062: 106:25//MOUSE MAMMARY TUMOR VIRUS (STRAIN BR6).//P10259

F-PLACE1004804

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F-PLACE1004813/HYPOTHETICAL PROTEIN UL12.//1.0:22:40//HUMAN CYTOMEGALOVIRUS (STRAIN 15 AD169).//P16777

F-PLACE1004814//HYPOTHETICAL 37.0 KD PROTEIN B0495.8 IN CHROMOSOME II.//2.8e-06:136:25// CAENORHABDITIS ELEGANS.//Q09217

F-PLACE1004815

F-PLACE1004824//HYPOTHETICAL 106.7 KD PROTEIN IN MUP1-SPR3 INTERGENIC REGION.//2.3e-09:70: 38//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53236

F-PLACE1004827//HYPOTHETICAL 9.4 KD PROTEIN IN FLAL 3'REGION (ORF3).//0.54:25:56//BACILLUS LI-CHENIFORMIS.//P22754

F-PLACE1004836//ANNEXIN VII (SYNEXIN) (FRAGMENT).//0.0066:12:66//BOS TAURUS (BOVINE).//P20072

25 F-PLACE1004840

F-PLACE1004868//MALE STERILITY PROTEIN 2.//4.0e-16:172:30//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//Q08891
F-PLACE1004885

F-PLACE1004900//MAST CELL DEGRANULATING PEPTIDE PRECURSOR (MCDP) (MCD) (PEPTIDE 401).//

90 1.0:23:47//APIS MELLIFERA (HONEYBEE).//P01499
F-PLACE1004902//PUTATIVE PRE-MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE

SPAC10F6.02C.//7.3e-15:94:47//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O42643
F-PLACE1004913//HYPOTHETICAL 7.2 KD PROTEIN IN BCSA-DEGR INTERGENIC REGION.//1.0:42:33//BA-

CILLUS SUBTILIS.//P54165
F-PLACE1004918//HYPOTHETICAL 12.4 KD PROTEIN IN RPS21B-MRS3 INTERGENIC REGION.//0.98:50:34//

SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P947012

F-PLACE1094930//HYPOTHETICAL PROTEIN MJ0562 //0.82-44:36//METHANOCOCCUS JANNASCHI.//

Q57982

F-PLACE1004934

40 F-PLACE1004937//HYPOTHETICAL 67.1 KD TRP-ASP REPEATS CONTAINING PROTEIN C57A10.05C IN CHROMOSOME I.//9.0e-10:87:33//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P87053

F-PLACE1004969//HYPOTHETICAL 55.1 KD PROTEIN B0416.5 IN CHROMOSOME X.//4.0e-14:184:25// CAENORHABDITIS ELEGANS.//Q11073

F-PLACE1004972//BROMELAIN INHIBITOR 2 (BI-II) (BROMELAIN INHIBITOR VI) (BI-VI).//1.0:35:37//ANANAS COMOSUS (PINEAPPLE).//P27478

F-PLACE1004979//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//5.3e-30:55:72//HOMO SAPIENS (HUMAN).// P39192

F-PLACE1004982//M PROTEIN, SEROTYPE 12 PRECURSOR (FRAGMENT).//0.00049:124:27//STREPTO-COCCUS PYOGENES.//P19401

F-PLACE1004985//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:26:34//LUMBRICUS TERRESTRIS (COMMON EARTHWORM).//Q34942

F-PLACE1005026//TELOMERE-BINDING PROTEIN HOMOLOG.//0.0011:179:27//EUPLOTES CRASSUS.// Q06183

F-PLACE1005027

55 F-PLACE1005046/JANNEXIN VII (SYNEXIN) (FRAGMENT).//0.082:44:36//BOS TAURUS (BOVINE).//P20072 F-PLACE1005052//MALE SPECIFIC SPERM PROTEIN MST84DD.//0.38:36:44//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//001645

F-PLACE1005055

- F-PLACE1005066//RING CANAL PROTEIN (KELCH PROTEIN).//2.9e-38:194:39//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- E-PLACE 1005077
- F-PLACE1005085//INSECT TOXIN 1 (BOT IT1).//0.85:36:33//BUTHUS OCCITANUS TUNETANUS (COMMON FUROPEAN SCORPION) //P55902
- F-PLACE1005086//!!!! ALÚ SUBFAMILY SQ WARNING ENTRY !!!!//8.5e-38:93:76//HOMO SAPIENS (HUMAN).// P39194
  - F-PLACE1005101//HYPOTHETICAL PROTEIN ZAP128 (FRAGMENT).//1.6e-11:35:100//HOMO SAPIENS (HU-MAN).//P49753
- F-PLACE1005102/IZINC FINGER PROTEIN 151 (POLYOMAVIRUS LATE INITIATOR PROMOTER BINDING PROTEIN) (IL-P1, IQINC FINGER PROTEIN 213), 30, e41-4110-38/MUS MUSCULUS (MOUSE),/060821 F-PLACE1005108/METALLOTHIONEIN-III (MT-III) (GROWTH INHIBITORY FACTOR) (GIF),//0.41.35:34/BOS TAURUS (BOVINE) / 197359
- F-PLACE 1005111//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L) (CHARGERIN II).//1.0:29:41//RATTUS NORVEGICUS (RAT).//P11608
- F-PLACE1005128//RABPHILIN-3A (FRAGMENT).//5.9e-05:95:36//MUS MUSCULUS (MOUSE).//P47708
  - F-PLACE1005146//FIBROBLAST GROWTH FACTOR INDUCIBLE PROTEIN 15 (FIN15).//0.17:48:35//MUS MUSCULUS (MOUSE).//Q61075
  - F-PLACE1005162//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.0e-31:60:76//HOMO SAPIENS (HUMAN).// P39189
  - E-PLACE1005176

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- F-PLACE1005181//HYPOTHETICAL 7 KD PROTEIN.//1.0:31:45//MEASLES VIRUS (STRAIN HALLE) (SUBA-CUTE SCLEROSE PANENCEPHALITIS VIRUS).//P06831
- F-PLACE1005187//GLUCAN SYNTHASE-1 (EC 2.4.1.34) (1,3-BETA-GLUCAN SYNTHASE) (UDP-GLUCOSE-
- 25 1,3-BETA-D-GLUCAN GLUCOSYLTRANSFERASE).//0.0025:58:34//NEUROSPORA CRASSA.//P38678 F-PLACE1005208//HYPOTHETICAL 10.7 KD PROTEIN.//0.34:57.42//VACCINIA VIRUS (STRAIN COPENHA-GEN)//P20511
  - F-PLACE1005232//AMELOGENIN, Y ISOFORM PRECURSOR.//0.70:60:35//HOMO SAPIENS (HUMAN).//
    Q99218
- 30 F-PLACE1005243//SERINE/THREONINE PROTEIN KINASE PKPA (EC 2.7.1.-).//0.0017:114:27//PHYCOMY-CES BLAKESLEEANUS.//Q01577
  - F-PLACE1005261//HYPOTHETICAL 90.8 KD PROTEIN T05H10.7 IN CHROMOSOME II.//1.2e-38:206:41// CAENORHABDITIS ELEGANS.//Q10003
  - F-PLACE1005266
- 35 F-PLACE1005277//PROTEIN GURKEN PRECURSOR.//0.58:95:29//DROSOPHILA MELANOGASTER (FRUIT FLY).//P42287
  - F-PLACE1005287//INNER CENTROMERE PROTEIN (INCENP).//2.0e-12:211:29//GALLUS GALLUS (CHICK-EN).//P53352
- F-PLACE1005305//GTP:AMP PHOSPHOTRANSFERASE MITOCHONDRIAL (EC 2.7.4.10) (AK3).//1.8e-78:205: 78//BOS TAURUS (BOVINE).//P08760
- F-PLACE1005308/WOUND-INDUCED BASIC PROTEIN.//0.99:40:40//PHASEOLUS VULGARIS (KIDNEY BEAN) (FRENCH BEAN).//0.09020
  - F-PLACE1005313//HYPOTHETICAL 8.7 KD PROTEIN IN LEUX-FECE INTERGENIC REGION (067).//0.15:36: 41//FSCHERICHIA COLI //P39355
- 45 F-PLACE1005327//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//1.0:19:52//HOMO SAPIENS (HUMAN)//P30808
  - F-PLACE1005331//BREAKPOINT CLUSTER REGION PROTEIN.//0.00021:98:35//HOMO SAPIENS (HUMAN).//
  - F-PLACE1005335//IROQUOIS-CLASS HOMEODOMAIN PROTEIN IRX-3.//0.37:98:33//MUS MUSCULUS (MOUSE).//P81067
  - F-PLACE1095373/PSEUDOURIDYLATE SYNTHASE 4 (EC.4.2.1.70) (PSEUDOURIDINE SYNTHASE 4) (TRNA PSEUDOURIDINE 55 SYNTHASE) (PSIS SYNTHASE) (PSIS SYNTHASE) (PSIS PSEUDOURIDYLATE SYNTHASE) (URACIL HYDROL-YASE) //0 010-99:28/ISACCHAROMYCES CEREVISIAE (BAKER'S YEAST) //P48567.
- F-PLACE1005374

- 55 F-PLACE1005409
  - F-PLACE1005453//LICHENASE PRECURSOR (EC 3.2.1.73) (ENDO-BETA-1,3-1,4 GLUCANASE).//1.0:50:32// NICOTIANA PLUMBAGINIFOLIA (LEADWORT-LEAVED TOBACCO).//P07979
  - F-PLACE1005467//KERATIN, FEATHER (F-KER).//0.0095:42:35//LARUS NOVAE-HOLLANDIAE (SILVER

### GULL) //P02451

- F-PLACE1005471//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3).//0.23:49:32//PHYTOPH-THORA INFESTANS (POTATO LATE BLIGHT FUNGUS).//Q37598
- F-PLACE1005477//HYPOTHETICAL PROTEIN ORF-1137.//9.6e-13:115:38//MUS MUSCULUS (MOUSE).// P11260
  - F-PLACE1005480//C-HORDEIN (CLONE PC HOR1-3) (FRAGMENT).//0.97:33:30//HORDEUM VULGARE (BAR-LFY) //P17991
  - F-PLACE1005481//HUNCHBACK PROTEIN (FRAGMENT).//0.30:52:38//APIS MELLIFERA (HONEYBEE).// P31504
- 10 F-PLACE1005494/TRANSIENT-RECEPTOR-POTENTIAL PROTEIN.//3.9e-05:87:33//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P19334

#### F-PLACE1005502

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- F-PLACE1005526//IMMEDIATE-EARLY PROTEIN IE180.//4.6e-05:132:32//PSEUDORABIES VIRUS (STRAIN KAPLAN) (PRV).//P33479
- 15 F-PLACE1005528//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//3.4e-09:31:74//HOMO SAPIENS (HUMAN).// P39195
  - F-PLACE1005530//HYPOTHETICAL 47.6 KD PROTEIN C16C10.5 IN CHROMOSOME III.//9.7e-50:148:58// CAENORHABDITIS ELEGANS.//Q09251
  - F-PLACE1005550//HYPOTHETICAL 40.2 KD PROTEIN K12H4.3 IN CHROMOSOME III.//3.0e-21:127:37// CAENORHABDITIS ELEGANS.//P34524
  - F-PLACE1005554//CYTOCHROME B (EC 1.10.2.2) (FRAGMENT).//0.84:38:31//DIPODOMYS CALIFORNICUS (KANGAROO RAT).//P16359
    - F-PLACE1005557//60S RIBOSOMAL PROTEIN L27. //4.8e-09:60:48//CRYPTOCOCCUS NEOFORMANS (FILO-BASIDIELLA NEOFORMANS). //P46288
- 25 F-PLACE1005574//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.89:44:29//BOS TAURUS (BOVINE).// P03929
  - F-PLACE1005584//MALE SPECIFIC SPERM PROTEIN MST87F.//0.00030:33:48//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P08175
- F-PLACE1005595//MMEDIATE-EARLY PROTEIN IE180.//0.00048:162:30//PSEUDORABIES VIRUS (STRAIN INDIANA-FUNKHAUSER / BECKER) (PRV).//P11675
  - F-PLACE1005603//HIGH-MOBILITY-GROUP PROTEIN (NONHISTONE CHROMOSOMAL PROTEIN).//
    0.00034:83:30//TETRAHYMENA PYRIFORMIS.//P40625
    - F-PLACE1005611//DNAJ PROTEIN.//8.6e-20:108:48//CLOSTRIDIUM ACETOBUTYLICUM.//P30725
- F-PLACE1005623//EXTRACELLULAR SIGNAL-REGULATED KINASE 5 (EC 2.7.1.-) (ERK5) (ERK4) (BMK1 KINASE).//0.80:116:31//HOMO SAPIENS (HUMAN).//Q13164
  - F-PLACE1005630//INTERLEUKIN-14 PRECURSOR (IL-14) (HIGH MOLECULAR WEIGHT B-CELL GROWTH FACTOR) (HMW-BCGF).//0.0024:74:39//HOMO SAPIENS (HUMAN).//P40222
  - F-PLACE 1005639/EXTRACELLULAR MATRIX PROTEIN 1 (SECRÉTORY COMPONENT P85) (FRAGMENT).//
    0.72:18:61/FRATTUS NORVEGICUS (RAT).//G022894
    F-PLACE 1005646/FINA HELICASF-LIKE PROTEIN DB10 //4 88-29:172:45//NICOTIANA SYLVESTRIS (WOOD
- TOBACCO).//P46942
  F-PLACE1005656//RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE M2 CHAIN (EC 1.17.4.1) (RIBONUCLE-
  - OTIDE REDUCTASE).//3.7e-64:133:75//MESOCRICETUS AURATUS (GOLDEN HAMSTER).//Q60561
    F-PLACE1005666//CHLOROPLAST 50S RIBOSOMAL PROTEIN L28.//0.57:36.41//PORPHYRA PURPUREA.//
- F-PLACE1005666//CHLOROPLAST 50S RIBOSOMAL PROTEIN L28.//0.57:36.41//PORPHYRA PURPUREA.// F-PLACE1005698/HYPOTHETICAL PROTEIN IN SIGD 3'REGION (ORFC) (FRAGMENT).//0.50:61:29//BACIL-
  - LUS SUBTILIS./IP40405
    F-PLACE1005727//ANTER-SPECIFIC PROLINE-RICH PROTEIN APG (PROTEIN CEX) (FRAGMENT).//0.46:
  - 27:51/BRASSICA NAPUS (RAPE).//P40603
    F-PLACE1005730//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENT).//0.95:21:52//ORYCTOLAGUS CUNICULUS
    - (RABBIT).//P02456
      F-PLACE1005739//INTERFERON-GAMMA INDUCIBLE PROTEIN MG11.//3.4e-46:111:53//MUS MUSCULUS
    - (MOUSE).//Q60710 F-PLACE1005755//HYPOTHETICAL 70.2 KD PROTEIN IN GSH1-CHS6 INTERGENIC REGION.//2.6e-12:66:
- 55 51//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P42951
  - F-PLACE1005763//S-ACYL FATTY ACID SYNTHASE THIOESTERASE, MEDIUM CHAIN (EC 3.1.2.14) (THIOESTERASE II).//1.5e-26:69:57//RATTUS NORVEGICUS (RAT).//P08635
    - F-PLACE1005799//KERATIN, ULTRA HIGH-SULFUR MATRIX PROTEIN (UHS KERATIN).//0.028:96:32//HOMO

- SAPIENS (HUMAN),//P26371
- F-PLACE1005802//PROTEIN PROSPERO.//0.86:64:42//DROSOPHILA MELANOGASTER (FRUIT FLY).//
- F-PLACE1005803//MYELOID DIFFERENTIATION PRIMARY RESPONSE PROTEIN MYD116.//1.0:95:25//MUS MUSCULUS (MOUSE).//P17564
- F-PLACE1005804//PROCESSING ALPHA-1,2-MANNOSIDASE (EC 3.2.1.-) (ALPHA-1,2-MANNOSIDASE 1B).//
  2.8e-73:198:73//MUS MUSCULUS (MOUSE) //P39098
  - F-PLACE1005813//HYPOTHETICAL 49.0 KD PROTEIN IN NSP1-KAR2 INTERGENIC REGION.//0.022:78:38// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47057
- F-PLACE1005828//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.8e-23:56:76//HOMO SAPIENS (HUMAN).// P39195
  - F-PLACE1005834//LATE CONTROL GENE B PROTEIN (GPB).//0.97:33:39//BACTERIOPHAGE 186.//P08711 F-PLACE1005845
  - F-PLACE1005850//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//5.5e-28:96:73//HOMO SAPIENS (HUMAN).// P39194
- 15 P39194 F-PLACE1005851

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- F-PLACE1005876//CLEAVAGE AND POLYADENYLATION SPECIFICITY FACTOR, 100 KD SUBUNIT (CPSF 100 KD SUBUNIT).//2.2e-99:155:95//BOS TAURUS (BOVINE).//Q10568
- 20 F-PLACE1005890//BEM46 PROTEIN (FRAGMENT).//1.8e-33:137:49//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P54069
  - F-PLACE1005898//NADH-UBIQUINONE OXIDOREDUCTASE MLRQ SUBUNIT (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-MLRQ) (CI-MLRQ).//0.77:58:34//HOMO SAPIENS (HUMAN).//000483
- F-PLACE1005921//AIG1 PROTEIN.//1.4e-23:165:38//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//
  P54120
- F-PLACE1005923//HYPOTHETICAL 22.4 KD PROTEIN (ORF16).//0.90:118:28//PARAMECIUM TETRAURE-
  - F-PLACE1005925//HYPOTHETICAL GENE 30 PROTEIN.//0.94:57:29//HERPESVIRUS SAIMIRI (STRAIN 11).//
- 30 F-PLACE1005932//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).// 0.42:128:32//NICOTIANA TABACUM (COMMON TOBACCO).//P13983
  - F-PLACE1005934//DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT (EC 2.7.7.6)(RPB1) (FRAG-MENT).//0.40:76:35//CRICETULUS GRISEUS (CHINESE HAMSTER).//P11414
  - F-PLACE1005936/TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN).//0.50:15:66//HUMAN IMMU-NODEFICIENCY VIRUS TYPE 1 (CLONE 12) (HIV-1).//P04326
    - F-PLACE1005951//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).// 0.0025:135:32//NICOTIANA TABACUM (COMMON TOBACCO).//P13983
    - F-PLACE1005953//HIGH POTENTIAL RON-SULFUR PROTEIN (HIPIP).//0.64:57:33//RHODOFERAX FER-MENTANS.//P80882
- 40 F-PLACE1005955//HYPOTHETICAL 54.2 KD PROTEIN IN ERP5-ORC6 INTERGENIC REGION.//1.0e-32:110: 50//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38821
  - F-PLACE1005966/TACHYPLESIN II PRECURSOR.//0.97:31:35//TACHYPLEUS TRIDENTATUS (JAPANESE HORSESHOE CRAB).//P14214
  - F-PLACE1005968//GATA FACTOR SREP.//0.17:52:40//PENICILLIUM CHRYSOGENUM.//Q92259
- 45 F-PLACE1005990//CELL PATTERN FORMATION-ASSOCIATED PROTEIN.//0.36:55:36//EMERICELLA NIDU-LANS (ASPERGILLUS NIDULANS).//P36011
  - F-PLACE1006002///!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.5e-36:102:75//HOMO SAPIENS (HU-MAN)//P39192
    F-PLACE1006003//HYPOTHETICAL 6.8 KD PROTEIN IN COX3-NAD1 INTERGENIC REGION (ORF 61).//1.0:
  - F-PLACE1006003//HYPOTHETICAL 6.8 KD PROTEIN IN COX3-NAD1 INTERGENIC REGION (ORF 61).//1.0 22:40//MARCHANTIA POLYMORPHA (LIVERWORT).//P38473
  - F-PLACE1086011/POLY [ADP-RIBOSE] POLYMERASE (EC 2.4.2.30) (PARP) (ADPRT) (NAD(+) ADP-RIBO-SYLTRANSFERASE) (POLY[ADP-RIBOSE] SYNTHETASE).//2.8e-21:163:36//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//0.11207
  - F-PLACE1006017//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.1e-10:43:67//HOMO SAPIENS (HUMAN).//
  - F-PLACE1006037/V/TELLOGENIN I PRECURSOR (VTG I) [CONTAINS: LIPOVITELLIN 1 (LV1); PHOSVITIN (PV); LIPOVITELLIN 2 (LV2)].//0.00019:123:37//FUNDULUS HETEROCLITUS (KILLIFISH) (MUMMICHOG).// 090508

- F-PLACE1006040//CAMP-REGULATED PHOSPHOPROTEIN 19 (ARPP-19).//3.2e-40:110:76//HOMO SAPIENS (HUMAN) //P56211
- F-PLACE1006076//BOWMAN-BIRK TYPE PROTEINASE INHIBITOR A-II.//0.99:30:40//ARACHIS HYPOGAEA (PEANUT).//P01066
- F-PLACE1006119//IMPORTIN BETA-3 SUBUNIT (KARYOPHERIN BETA-3 SUBUNIT) (RAN-BINDING PROTEIN 5).//8.8e-94:218:76//HOMO SAPIENS (HUMAN).//000410
  - F-PLACE1006129/IA-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//0.00092:228:26//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323
  - F-PLACE1006139//HYPOTHETICAL 52.9 KD PROTEIN IN SAP155-YMR31 INTERGENIC REGION.//5.9e-55:
  - 128:50//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P43616
    F-PLACE1006143///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.4e-25:107:63//HOMO SAPIENS (HU-MAN).//P39194
  - F-PLACE1008157/IE-SELECTIN PRECURSOR (ENDOTHELIAL LEUKOCYTE ADHESION MOLECULE 1) (ELAM-1) (LEUKOCYTE-ENDOTHELIAL CELL ADHESION MOLECULE 2) (LECAM2) (CD62E).//1.3e-21:168: 32//SUS SCROFA (PIG).//PS8110
  - F-PLACE1006159//COLD SHOCK INDUCED PROTEIN TIR1 PRECURSOR (SERINE-RICH PROTEIN 1).//0.46: 98:29//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P10863
    - F-PLACE1006164//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3) //0:70:28:42//ARTEMIA SALINA (BRINE SHRIMP) //P19049
- F-PLACE1006167//GLUCÓAMYLASE S1/S2 PRECURSOR (EC 3.2.1.3) (GLUCAN 1,4-ALPHA- GLUCOSI-DASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//8.9e-05:167:32//SACCHAROMYCES CEREVISIAE (BAKERS YEAST)/P00840
- F-PLACE10817/JALPHA-ADAPTIN C (CLATHRIN ASSEMBLY PROTEIN COMPLEX 2 ALPHA-C LARGE CHAIN) (100 KD COATED VESICLE PROTEIN C) (PLASMA MEMBRANE ADAPTOR HA2/AP2 ADAPTIN ALPHA
- 25 C SUBUNIT).//1 1e-67:157:88//MUS MUSCULUS (MOUSE).//P17427 F-PLACE1006187//G1/S-SPECIFIC CYCLIN E.//5.6e-75:224:62//HOMO SAPIENS (HUMAN).//P24864
  - F-PLACE1006195/T-RELATED PROTEIN (TRP) (BRACHYENTERON PROTEIN).//0.99:177:29//DROSOPHILA MELANOGASTER (FRUIT FLY).//P55965
  - F-PLACE1006196//PUTATIVE ATP-DEPENDENT RNA HELICASE C12C2.06.//2.0e-33:183:46//SCHIZOSAC-CHAROMYCES POMBE (FISSION YEAST).//Q09747
  - F-PLACE1006205

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- F-PLACE1006223//SPERM MITOCHONDRIAL CAPSULE SELENOPROTEIN (MCS).//0.00015:22:50//MUS MUSCULUS (MOUSE).//P15265
- F-PLACE1006225//VIRION INFECTIVITY FACTOR (SOR PROTEIN).//1.0:63:34//HUMAN IMMUNODEFICIEN-CY VIRUS TYPE 1 (NDK ISOLATE) (HIV-1).//P18805
- F-PLACE1006236
  - F-PLACE1006239//60S ACIDIC RIBOSOMAL PROTEIN P2 (FRAGMENT).//0.48:23:52//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//P51407
- F-PLACE1006246//CMP-SIALIC ACID TRANSPORTER (CMP-SIA-TR).//0.012:84:30//MUS MUSCULUS (MOUSE).//Q61420
  - F-PLACE1006248//140 KD NUCLEOLAR PHOSPHOPROTEIN (NOPP140).//0.017:203:22//RATTUS NORVEGI-CUS (RAT).//P41777
    - F-PLACE1006262//L-FUCULOSE PHOSPHATE ALDOLASE (EC 4.1.2.17).//0.84:25:52//HAEMOPHILUS INFLU-FNZAF //P44777
- 45 F-PLACE1006288
  - F-PLACE1006318//CYSTEINE-RICH ANTIFUNGAL PROTEIN 1 (AFP1) (M1).//1.0:29:48//SINAPIS ALBA (WHITE MUSTARD) (BRASSICA HIRTA).//P30231
  - F-PLACE1006325//CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT KINASE INHIBITOR P57) (P57KIP2).//0.99:97:32//HOMO SAPIENS (HUMAN).//P49918
  - F-PLACE1006335//PROLINE-RICH PEPTIDE P-B.//0.56:19:52//HOMO SAPIENS (HUMAN).//P02814
  - F-PLACE1006357
    - F-PLACE1006360
    - F-PLACE1006368//NUF1 PROTEIN (SPINDLE POLY BODY SPACER PROTEIN SPC110),//0.0057:122:31//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST),//P32380
- 55 F-PLACE1006371//ARS BINDING PROTEIN 1.//0.00030:142:30//SCHIZOSACCHAROMYCES POMBE (FIS-SION YEAST).//P49777
  - F-PLACE1006382//NEUROTOXIN V.//0.85:28:39//ANDROCTONUS MAURETANICUS MAURETANICUS (SCORPION).//P01482

- F-PLACE1006385//HYPOTHETICAL 45.1 KD PROTEIN IN RPS5-ZMS1 INTERGENIC REGION.//3.1e-35:165: 47//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47160
- F-PLACE1006412//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//4.3e-08:40:47//HOMO SAPIENS (HU-MAN).//P08547
- 5 F-PLACE1006414//FORKHEAD-RELATED TRANSCRIPTION FACTOR 4 (FREAC-4).//3.8e-05:123:39//HOMO SAPIENS (HUMAN).//Q16676
  - F-PLACE1006438//ZINC FINGER PROTEIN 165.//2.8e-21:76:64//HOMO SAPIENS (HUMAN).//P49910
  - F-PLACE1006445//SUPPRESSOR OF HAIRY WING PROTEIN.//0.058:99:29//DROSOPHILA VIRILIS (FRUIT FLY).//Q08876
- F-PLACE1006469//ACETYL-COENZYME A SYNTHETASE (EC 6.2.1.1) (ACETATE--COA LIGASE) (ACYL- AC-TIVATING ENZYME),//1.8e-64:177:50//ESCHERICHIA COLI,//P27550
  - F-PLACE1006470
  - F-PLACE1006482//TRANSCRIPTION FACTOR MAFF.//2.0e-47:120:85//GALLUS GALLUS (CHICKEN).//
- 15 F-PLACE1006488//SIGNAL RECOGNITION PARTICLE 68 KD PROTEIN (SRP68).//1.8e-85:173:95//CANIS FA-MILIARIS (DOG).//000004
  - F-PLACE1006492/VERY HYPOTHETICAL 11.2 KD PROTEIN C56F8.13 IN CHROMOSOME I.//0.75:32:56// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10261
  - F-PLACE1006506 F-PLACE1006521

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- F-PLACE1006531//HYPOTHETICAL 115.4 KD PROTEIN ZK757.3 IN CHROMOSOME III.//1.3e-53:167:61// CAENORHABDITIS ELEGANS.//P34681
- F-PLACE1006534
- F-PLACE1006540
- F-PLACE1006552//MYOSIN HEAVY CHAIN, CLONE 203 (FRAGMENT).//1.3e-07:242:23//HYDRA ATTENUATA (HYDRA) (HYDRA VULGARIS).//P39922
  - F-PLACE1006598///!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!!//0.17:43:51//HOMO SAPIENS (HUMAN).// P39190
- F-PLACE1006615//ACROSIN PRECURSOR (EC 3.4.21.10).//3.6e-05:66:43//ORYCTOLAGUS CUNICULUS (RABBIT).//P48038
- F-PLACE1006617//HYPOTHETICAL 14.6 KD PROTEIN (READING FRAME C) (REPUCATION).//1.0:74:29// STAPHYLOCOCCUS AUREUS //P03861
  - F-PLACE1006626//HYPOTHETICAL HELICASE K12H4.8IN CHROMOSOME III.//2.9e-10:73:46//CAENORHAB-DITIS ELEGANS.//P34529
- 35 F-PLACE1006629//HYPOTHETICAL PROTEIN BB0410.//1.0:23:43//BORRELIA BURGDORFERI (LYME DIS-EASE SPIROCHETE).//051371
  - F-PLACE1006640
  - F-PLACE1006673
  - F-PLACE1006678//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENT).//1.0:36:41//ORYCTOLAGUS CUNICULUS (RABBIT).//P02456
- F-PLACE1006704//BROAD-COMPLEX CORE-TNT1-Q1-Z1 PROTEIN (BRCORE-TNT1-Q1-Z1) [CONTAINS: BROAD-COMPLEX CORE-Q1-Z1
  - PROTEINI.//0.00062:157:26//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q01295
- F-PLACE 1006731//RIBOFLAVIN KINASE (EC 2.7.1.26) (FLAVOKINASE) / FMN ADENYLYLTRANSFERASE (EC 2.7.7.2) (FAD PYROPHOSPHORYLASE) / FAD SYNTHETASE) //1.3e-07:127:36//CORYNEBACTERIUM AMMONIAGENES ///658263
  - F-PLACE1006754//CARCINOEMBRYONIC ANTIGEN CGM1 PRECURSOR (CD66D ANTIGEN).//1.9e-19:78: 53//HOMO SAPIENS (HUMAN).//P40198
- F-PLACE1006760//COLLAGEN ALPHA 1(III) CHAIN (FRAGMENT).//0.21:107:30//RATTUS NORVEGICUS (RAT).//P13941
  - F-PLACE 1006779//CYTOTOXIN 5 (CTX V).//1.0:20:30/INAJA MOSSAMBICA (MOZAMBIQUE COBRA)./IP25517 F-PLACE 1006782//ZINC FINGER PROTEIN 1.//0.00052:178:28//CANDIDA ALBICANS (YEAST)./IP28875 F-PI ACE 1006792
- F-PLACE1006795//VOLTAGE-GATED POTASSIUM CHANNEL PROTEIN SHAW (SHAW2).//1:0:80:30//DRO-SOPHILA MELANOGASTER (FRUIT FLY).//P17972
- F-PLACE1006800//HYPOTHETICAL 9.4 KD PROTEIN.//0.99:62:33//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20569
  - F-PLACE1006805

- F-PLACE1006815//HYPOTHETICAL PROTEIN UL61.//0.038:146:32//HUMAN CYTOMEGALOVIRUS (STRAIN AD1691 //P16818
- F-PLACE1006819//LINE-1 REVERSE TRANSCRIPTASE HOMOLOGJ/7.3e-98:239:76//HOMO SAPIENS (HU-MAN).//P08547
- F-PLACE100829/UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 8 (EC 3.1.2.15) (UBIQUITIN THIOLESTE-RASE 8), UBIQUITIN PSPECIFIC PROCESSIND PROTEASE 8) (DEUBIQUITINATING ENZYME 8),//0.081:34: 58//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P50102 F-PLACE1008801
  - F-PLACE 1006860
- F-PLACE1006878//HYPOTHETICAL 8.2 KD PROTEIN IN MOBL 3'REGION (ORF 3).//0.85:27:37//THIOBACIL-LUS FERRODXIDANS //P20087
  - F-PLACE1006883//VITAMIN D3 RECEPTOR (VDR) (1,25-DIHYDROXYVITAMIN D3 RECEPTOR).//0.78:51:37// MUS MUSCULUS (MOUSE).//P48281
- F-PLACE1006901/HYPOTHETICAL 8.1 KD PROTEIN.//0.99:55:23//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20567
- F-PLACE1006904//MATING-TYPE LOCUS ALLELE B1 PROTEIN://0.95:86:26//USTILAGO MAYDIS (SMUT FUNCIJS)//P22015
  - F-PLACE 1006917//HYPOTHETICAL 40.9 KD PROTEIN C08B11.5 IN CHROMOSOME II.//6.9e-15:101:45//
    CAENORHABDITIS ELEGANS //009442
- 20 F-PLACE1006932/HISTIDINE-RICH, METAL BINDING POLYPEPTIDE://0.089:28:39/HELICOBACTER PY-LORI (CAMPYLOBACTER PYLORI)//048251
  - F-PLACE1006935/HYPOTHETICAL 95.2 KD PROTEIN R144.6 IN CHROMOSOME III.//0.93:35:48// CAENORHABDITIS ELEGANS.//Q10000
- F-PLACE1006956/TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII135)

  25 (TAFII-130) (TAFII130) //0.00079-122-36/HOMO SAPIENS (HUMAN) //0.00268
  - F-PLACE1006958//OSMOTIC STRESS PROTEIN 94 (HEAT SHOCK 70-RELATED PROTEIN APG-1).//8.8e-70: 140-98//MUS UNUS (MOUSE).//P48722 F-PLACE1006961
  - F-PLACE1006962/IAPOLIPOPROTEIN C-I PRECURSOR (APO-C1).//1.0:25:40//PAPIO HAMADRYAS (HAMAD-RYAS BABOON).//P34929
- 39 RYAS BABOON),//P34929 F-PLACE1006966//HYPOTHETICAL 49.1 KD PROTEIN IN SSB2-SPX18 INTERGENIC REGION.//1.6e-47:221: 45//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P40160
  - F-PLACE1006989//HYPOTHETICAL 13.1 KD HIT-LIKE PROTEIN IN P37 5'REGION.//0.15:46:32//MYCOPLAS-MA HYORHINIS.//P32083
- 35 F-PLACE1007014//36 KD NUCLEOLAR PROTEIN HNP36 (DELAYED-EARLY RESPONSE PROTEIN 12) (DER12).//3.4e-09:120:29//HOMO SAPIENS (HUMAN).//Q14542
  - F-PLACE1007021//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00046:42:59//HOMO SAPIENS (HUMAN).// P39188
- F-PLACE1007045//HYPOTHETICAL PROTEIN ORF-1137.//8.1e-14:115:35//MUS MUSCULUS (MOUSE).//
  40 P11260
  - F-PLACE1007053//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.038:48:39//HOMO SAPIENS (HUMAN).//P22531
  - F-PLACE1007068//PROTEIN-LYSINE 6-OXIDASE PRECURSOR (EC 1.4.3.13) (LYSYL OXIDASE).//0.0040:113: 39//GALLUS GALLUS (CHICKEN).//Q05063
- 45 F-PLACE1007097/HYPOTHETICAL 6.8 KD PROTEIN IN HE65-PK2 INTERGENIC REGION.//0.97:47:29// AUTOGRAPHA CALIFOTNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41663 F-PLACE1007105//HYPOTHETICAL 83.6 KD PROTEIN C15A10.10 IN CHROMOSOME L//2.9e-33:219:37//
  - F-PLACE1007105/HYPOTHETICAL 83.6 KD PROTEIN C15A10.10 IN CHROMOSOME L//2.9e-33:219: SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//013730 F-PLACE1007111
  - F-PLACE1007111
- 50 F-PLACE1007112//HYPOTHETICAL 9.2 KD PROTEIN.//0.47:75:28//ESCHERICHIA COLI.//P03853
  - F-PLACE1007132//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.8e-11:56:57//HOMO SAPIENS (HUMAN).// P39188
  - F-PLACE1007140//GAR2 PROTEIN.//0.72:185:24//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).// P41891
- 55 F-PLACE1007178//HYPOTHETICAL 8.5 KD PROTEIN CY274.40C.//0.97:79:30//MYCOBACTERIUM TUBER-CUI OSIS //Q10826
  - F-PLACE1007226/HYPOTHETICAL 42.6 KD PROTEIN IN GSHB-ANSB INTERGENIC REGION (0378).//1.9e-15:123:32//ESCHERICHIA COLI.//P52062

- F-PLACE1007238//MYOSIN HEAVY CHAIN IB (MYOSIN HEAVY CHAIN IL).//5.5e-10:98:44//ACANTHAMOEBA CASTELLANII (AMOEBA).//P19706
- F-PLACE1007239//TRANSCRIPTION ELONGATION FACTOR S-II (TRANSCRIPTION ELONGATION FACTOR A) //3.9e-19:96:57//HOMO SAPIENS (HUMAN) //P23193
- F-PLACE1007242//GUANINE NUCLEOTIDE DISSOCIATION STIMULATOR RALGDS FORM B (RALGEF).//1.0: 132:30//RATTUS NORVEGICUS (RAT).//Q03386
  - F-PLACE1007243/HyPOTHETICAL 53.3 KD PROTEIN IN HXT8-CAN1 INTERGENIC REGION.//0.041:114:29// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39981
- F-PLACE1007257//DIAPHANOUS PROTEIN.//1.3e-42:205:46//DROSOPHILA MELANOGASTER (FRUIT FLY)//P48608
  - F-PLACE1007274//CADMIUM-METALLOTHIONEIN (CD-MT).//0.054:60:30//HELIX POMATIA (ROMAN SNAIL) (EDIBLE SNAIL).//P33187
  - F-PLACE1007276//BETA-DEFENSIN 1 PRECURSOR (RHBD-1) (DEFENSIN, BETA 1).//1.0:42:28//SUS SCRO-FA (PIG1//O62697
- 15 F-PLACE1007282/JOUTER CAPSID PROTEIN VP4 (HEMAGGLUTININ) (OUTER LAYER PROTEIN VP4) [CONTAINS: OUTER CAPSID PROTEINS VP5 AND VP8].//0.070:126:27//HUMAN ROTAVIRUS (SEROTYPE 4 / STRAIN ST. THOMAS 3).//P11200
  - F-PLACE1007286

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- F-PLACE1007301//HYPOTHETICAL PROTEIN KIAA0168.//0.042:61:39//HOMO SAPIENS (HUMAN).//P50749 F-PLACE1007317
  - F-PLACE1007342//PROTEIN GRAINY-HEAD (DNA-BINDING PROTEIN ELF-1) (ELEMENT I-BINDING ACTIV-ITY) (TRANSCRIPTION FACTOR NTF-1).//1.7e-06:77:36//DROSOPHILA MELANOGASTER (FRUIT FLY).//
- F-PLACE1007346//TRANSCRIPTION INTERMEDIARY FACTOR 1-BETA (KRAB-A INTERACTING PROTEIN)

  (KRIP-1\/0.0026:147:27//MUS MUSCULUS (MOUSE\/062318
  - F-PLACE1007367//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.3e-37:110:76//HOMO SAPIENS (HU-MAN).//P39189
  - F-PLACE1007375//PHORBOL ESTER/DIACYLGLYCEROL-BINDING PROTEIN UNC-13.//4.7e-07:71:39//
    CAENORHABDITIS ELEGANS.//P27715
- 30 F-PLACE1007386/HYPOTHETICAL 7.6 KD PROTEIN IN FLO1-PHO11 INTERGENIC REGION.//0.74:48:29// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39561
  - F-PLACE1007402//TRANSCRIPTIONAL REGULATORY PROTEIN ENTR (ENTERICIDIN R).//0.99:63:36//CIT-ROBACTER FREUNDII.//O69280
    - F-PLACE1007409//WHITE PROTEIN.//7.9e-38:179:41//DROSOPHILA MELANOGASTER (FRUIT FLY).//
      - F-PLACE1007416/IDIPEPTIDYL PEPTIDASE IV (EC 3.4.14.5) (DPP IV) (T-CELL ACTIVATION ANTIGEN CD26) (TP103) (ADENOSINE DEAMINASE COMPLEXING PROTEIN-2) (ADABP).//0.031:159.23//HOMO SAPIENS (HUMAN)./PZ7487
- F-PLACE1007450//ZINC FINGER PROTEIN 39 (ZINC FINGER PROTEIN KOX27) (FRAGMENT).//0.023:36:50//
  HOMO SAPIENS (HUMAN).//P17038
  - F-PLACE1007452//HYPOTHETICAL 22.1 KD PROTEIN IN CCP1-MET1 INTERGENIC REGION.//2.2e-18:85: 54//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36149
    - F-PLACE1007454//PHOTOSYSTEM II REACTION CENTRE N PROTEIN.//0.66:13:53//CHLAMYDOMONAS RE-INHARDTII.//0.06480
- 45 F-PLACE1007460/IATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L)./JO 93:45:33//SUS SCROPA (PIG).//O3:5914 F-PLACE1007478//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASEJ/6.3e-08:50:56/MUS MUSCULUS (MOUSE)//P11369
  - F-PLACE1007484//HYPOTHETICAL 6.8 KD PROTEIN IN REPLICATION ORIGIN REGION.//0.87:43:37//ES-CHERICHIA COLI.//P03849
  - F-PLACE1007488//PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE FACTOR (RHO/RAC GEF) (FA-CIOGENITAL DYSPLASIA PROTEIN)//1.2e-25:202:31/HOMO SAPIENS (HUMAN) //P98174 F-PLACE1007507/HYPOTHETICAL 160 KD PROTEIN IN TAF60-64P1 INTERGENIC REGION.//0.12:128:25//
    - SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53139
- F-PLACE1007511//KERATIN, TYPE I CYTOSKELETAL 19 (CYTOKERATIN 19) (K19) (CK 19).//2.1e-45:209:48//
  BOS TAURUS (BOVINE).//P08728
  - F-PLACE1007524//HYPOTHETICAL 9.2 KD PROTEIN.//0.74:80:30//VACCINIA VIRUS (STRAIN COPENHA-GEN).//P20550
  - F-PLACE1007525

- F-PLACE1007537//MYOTROPHIN (V-1 PROTEIN) (GRANULE CELL DIFFERENTIATION PROTEIN).//0.045:92: 30//MUS MUSCULUS (MOUSE), AND RATTUS NORVEGICUS (RAT).//P80144
- F-PLACE1007544//IMMEDIATE-EARLY PROTEIN IE180.//1.5e-07:59:50//PSEUDORABIES VIRUS (STRAIN KAPLAN) (PRV).//P33479
- 5 F-PLACE1007547//HYPOTHETICAL 97.1 KD PROTEIN R05D3.4 IN CHROMOSOME III.//2.5e-16:188:34// CAENORHABDITIS ELEGANS.//P34537 F-PLACE1007557
  - F-PLACE1007583//PROLINE RICH 33 KD EXTENSIN-RELATED PROTEIN PRECURSOR (FRAGMENT).//0.98: 72:33//DAUCUS CAROTA (CARROT). //P06600
- F-PLACE1007598//ZINC FINGER PROTEIN 92 (ZINC FINGER PROTEIN HTF12) (FRAGMENT).//1.7e-11:88: 43//HOMO SAPIENS (HUMAN).//Q03936
  - F-PLACE1007618//ANION EXCHANGE PROTEIN 2 (NON-ERYTHROID BAND 3-LIKE PROTEIN) (B3RP).//0.19: 109:27//MUS MUSCULUS (MOUSE).//P13808
- F-PLACE1007621//PHOSPHATE REGULON SENSOR PROTEIN PHOR (EC 2.7.3.-) (FRAGMENT).//0.98:34: 41//PSEUDOMONAS AERUGINOSA.//P23621
- F-PLACE1007632//COLLAGEN ALPHA 2(I) CHAIN (FRAGMENT).//0.70:110:34//BOS TAURUS (BOVINE).//
  - F-PLACE1007645//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//0.99:20:45//STRUTHIO CAMELUS (OSTRICH).//021401
- 20 F-PLACE1007649//GLUCOAMYLASE S1/S2 PRECURSOR (EC 3.2.1.3) (GLUCAN 1,4-ALPHA- GLUCOSI-DASE) (1.4-ALPHA-D-GLUCAN GLUCOHYDROLASE).//8.1e-06:197:26//SACCHAROMYCES CEREVISIAE (BAKERS YEAST)/P00840
  - F-PLACE1007677//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//1.0:47:46//HOMO SAPIENS (HUMAN).//
- 25 F-PLACE1007688//LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA AUTOANTIGEN HOMOLOG).// 2.7e-06:116:28//AEDES ALBOPICTUS (FOREST DAY MOSQUITO).//Q26457
  - F-PLACE1007690//SPERM PROTAMINE P1.//0.12:26:50//TACHYGLOSSUS ACULEATUS ACULEATUS (AUSTRALIAN ECHIDNA).//P35311
- F-PLACE1007697/ISPERM PROTAMINE PI.//0.19.34.52//DIDELPHIS MARSUPIALIS VIRGINIANA (NORTH AMERICAN OPOSSUM), AND MONODELPHIS DOMESTICA (SHORT-TAILED GREY OPOSSUM).//P35305 F-PLACE1007705/IBIOH PROTEIN.//0.015.97.29/IESCHERICHIA COLI.//P13001
  - F-PLACE1007706//HYPOTHETICAL 112.2 KD PROTEIN IN TIF35-NPL3 INTERGENIC REGION (ORF1).//5.3e-55:190:56//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//F32898
- F-PLACE1007725

  5-5-PLACE1007729//PROTEASE (EC 3.4.23.-).//1.8e-21;136:42//MOUSE MAMMARY TUMOR VIRUS (STRAIN)
  - BR6).//P10271

    F-PLACE:1007730//SALIVARY PROLINE-RICH PROTEIN II-1 (FRAGMENT).//0.0031:77:40//HOMO SAPIENS
    (HIJMAN) //P81489
- F-PLACE1007737//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.78:39:56//HOMO SAPIENS (HUMAN).//
  40 P39195
- F-PLACE1007743

- F-PLACE1007746//RRP5 PROTEIN HOMOLOG (KIAA0185) (FRAGMENT).//0.0066:168:25//HOMO SAPIENS (HUMAN).//Q14690
- F-PLACE1007791/KRUEPPEL PROTEIN (FRAGMENT).//0.62:17:41/LITHOBIUS FORFICATUS.//Q01872 F-PLACE1007807//HYPOTHETICAL 6.4 KD PROTEIN IN BLTR-SPOIIIC INTERGENIC REGION.//1.0:40:30//BA-
- CILLUS SUBTILIS./IP54446
  F-PLACE1007810/IANTHOPLEURIN A (TOXIN AP-A).//0.79:28:46//ANTHOPLEURA XANTHOGRAMMICA (GI-ANT GREEN SEA ANEMONE).//P01530
  - F-PLACE1007829//SPORE COAT PROTEIN G.//1.0:65:38//BACILLUS SUBTILIS.//P39801
  - F-PLACE 1007 829/37 GRE COAT PROTEIN G.// 1.0.03.30//DAGIEEGS 30BTEIS://P39
  - F-PLACE1007846//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.5e-32:37:94//HOMO SAPIENS (HU-MAN).//P08547
    - F-PLACE1007852//RHO-RELATED GTP-BINDING PROTEIN RHOH (GTP-BINDING PROTEIN TTF).//8.7e-05: 138:30//HOMO SAPIENS (HUMAN).//Q15669
- 55 F-PLACE1007858//ANAPHASE SPINDLE ELONGATION PROTEIN.//0.0039:127:25//SACCHAROMYCES CER-EVISIAE (BAKER'S YEAST).//P50275
  - F-PLACE1007866
  - F-PLACE1007877

- F-PLACE 1007897//CD44 ANTIGEN PRECURSOR (PHAGOCYTIC GLYCOPROTEIN I) (PGP-1) (HUTCH-I) (EXTRACELLULAR MATRIX RECEPTOR-II) (ECMR-III) (PGP0 LYMPHOCYTE HOMING/ADHESION RECEPTOR) (HERMES ANTIGEN) (HYALURONATE RECEPTOR) (HEPARAN SULFATE PROTEOGLYCAN) (HAM1 ANTIGEN) // (AV128-28/JMESOCRICETUS AURATUS (GOLDEN HAMSTER) // (960522
- F-PLACE1007908//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//5.5e-28:61:65//HOMO SAPIENS (HUMAN).// P39192
  - F-PLACE1007946//HYPOTHETICAL MERCURIC RESISTANCE PROTEIN MERC.//0.84:48:37//PSEU-DOMONAS AERUGINOSA.//P04139
- F-PLACE1007954//HYPOTHETICAL 45.5 KD PROTEIN IN FIG1-GIP1 INTERGENIC REGION.//0.00070:96:29//
  98ACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38226
  - F-PLACE1007955//HYPOTHETICAL 84.3 KD PROTEIN ZK945.10 IN CHROMOSOME II.//0.00027:255:23//
    CAENORHABDITIS ELEGANS.//Q09625
  - F-PLACE1007958//HIGH-AFFINITY CAMP-SPECIFIC 3',5'-CYCLIC PHOSPHODIESTERASE (EC 3.1.4.17).//
    1.7e-09:127:30//MUS MUSCULUS (MOUSE).//P70453
- 15 F-PLACE1007969//HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION.//2.4e-05:104:37// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41479
  - F-PLACE1007990//SPERM PROTAMINE P1.//0.78:36:47//ORNITHORHYNCHUS ANATINUS (DUCKBILL PLAT-YPUS).//P35307
- F-PLACE1008000//CHANNEL ASSOCIATED PROTEIN OF SYNAPSE-110 (CHAPSYN-110) (SYNAPTIC DEN-20 SITY PROTEIN PSD-93).//1.2e-16:128:39//RATTUS NORVEGICUS (RAT).//Q63622
- F-PLACE1008002
  F-PLACE1008044/NUCLEAR PORE COMPLEX PROTEIN NUP107 (NUCLEOPORIN NUP107) (107 KD NU-
  - CLEOPORIN) (P105).//3.9e-106:208:93//RATTUS NORVEGICUS (RAT).//P52590
- F-PLACE1008045//COLLAGEN ALPHA 1(XII) CHAIN (FRAGMENTS).//3.9e-09:49:53//BOS TAURUS (BO-VINE).//P25508
  - F-PLACE1008080//RNA REPLICASE POLYPROTEIN (EC 2.7.7.48).//0.00025:100:27//EGGPLANT MOSAIC VI-RUS.//P20126
- F-PLACE1008095//PROTOPORPHYRINOGEN OXIDASE (EC 1.3.3.4) (PPO).//0.90:74:25//MYCOBACTERIUM TUBERCULOSIS.//053230
- F-PLACE1008111//HYPOTHETICAL PROTEIN MJECS12.//0.30:38:42//METHANOCOCCUS JANNASCHII.// Q60311
  - F-PLACE1008122//PEA2 PROTEIN (PPF2 PROTEIN).//0.0085:117:34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40091
  - F-PLACE1008129//PROCOLLAGEN ALPHA 2(I) CHAIN PRECURSOR (FRAGMENTS).//1.8e-06:154:36//GAL-LUS GALLUS (CHICKEN).//P02467
    - F-PLACE1008132//HYPOTHETICAL 127.4 KD PROTEIN F07F6.4 IN CHROMOSOME III.//1.4e-13:227:36// CAENORHABDITIS ELEGANS.//Q09531
    - F-PLACE1008177//TRICHOHYALIN.//2.7e-10:230:26//OVIS ARIES (SHEEP).//P22793
- F-PLACE1008181

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- 40 F-PLACE1008198//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//0.00044:121:34// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437
  - F-PLACE1008201//ZINC FINGER PROTEIN ZFMSA12A.//3.0e-05:82:37//MICROPTERUS SALMOIDES (LARGEMOUTH BASS).//P38621
- F-PLACE1008209//METALLOTHIONEIN-I (MT-I).//0.95:39:35//CERCOPITHECUS AETHIOPS (GREEN MON-KEY) (GRIVET).//P02797
  - F-PLACE1008231//PROCYCLIC FORM SPECIFIC POLYPEPTIDE B1-ALPHA PRECURSOR (PROCYCLIN) (PARP).//0.028:23:52//TRYPANOSOMA BRUCEI BRUCEI.//P08469 F-PLACE1008244//YEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//2.2e-23:148:38//PODOSPORA AN-INCECTION PROTEIN FROM THE PROTEIN FROM THE
  - F-PLACE1008244//VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1.//2.2e-23:148:38//PODOSPORA AN SERINA.//Q00808
  - F-PLACE1008273//COATOMER GAMMA SUBUNIT (GAMMA-COAT PROTEIN) (GAMMA-COP).//1.1e-97:222: 81/BOS TAURUS (BOVINE).//P53620
    - F-PLACE1008275//DNA REPAIR PROTEIN REV1 (EC 2.7.7.-).//5.8e-20:161:37//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P12689
- F-PLACE1008280//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.1e-23:124:42//HOMO SAPIENS (HU-55 MAN).//P08547
  - F-PLACE1008309/HYPOTHETICAL 98.3 KD PROTEIN C9G1.06C IN CHROMOSOME I.//0.47:99:37// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//014302
    - F-PLACE1008329//PUTATIVE Z PROTEIN.//0.73:52:28//OVIS ARIES (SHEEP).//P08105

- F-PLACE1008330//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/9.0e-37:75:81//HOMO SAPIENS (HUMAN).// P39194
- F-PLACE1008331//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.1e-08:70:50//HOMO SAPIENS (HUMAN).//
- F-PLACE1008356//FRUIT PROTEIN PKIWI501.//0.0037:148:29//ACTINIDIA CHINENSIS (KIWI) (YANGTAO).//
  - F-PLACE1008368/RING CANAL PROTEIN (KELCH PROTEIN).//3.5e-18:205:30//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652 F-PLACE1008369
- F-PLACE1008369
- 10 F-PLACE1008392
  - F-PLACE1008398//GENE 33 POLYPEPTIDE.//1.5e-102:225:84//RATTUS NORVEGICUS (RAT).//P05432
- F-PLACE1008401//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//2.9e-08:186:34//MUS MUSCULUS (MOUSE).//P05143
- F-PLACE1008402//GENERAL VESICULAR TRANSPORT FACTOR P115 (TRANSCYTOSIS ASSOCIATED PROTEIN) (TAP).//9.4e-105:207:98//BOS TAURUS (BOVINE).//P41541
- F-PLACE1008405
  - F-PLACE1008424/I/PROTEIN UL56.//1.0:65:33//HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN HFEM)./I/P36297
    F-PLACE1008426/I/MYOSIN HEAVY CHAIN, NON-MUSCLE (ZIPPER PROTEIN) (MYOSIN II)./I/4.4e-05:185:28//
    DROSOPHILA MEI ANOGASTER (FRUIT FLY) ///099323
- F-PLACE1008429/NEURONAL AXONAL MEMBRANE PROTEIN NAP-22.//0.00054:172:25//RATTUS NOR-VEGICUS (RAT).//Q05175
  - F-PLACE1008437//HYPOTHETICAL 115.4 KD PROTEIN ZK757.3 IN CHROMOSOME III.//1.9e-23:226:34//
- F-PLACE1008455//DNA-BINDING PROTEIN (AGNOPROTEIN).//0.97:23:52//BUDGERIGAR FLEDGLING DIS-25 EASE VIRUS (BFDV).//P13893
  - F-PLACE1008457//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.1e-12:89:47//HOMO SAPIENS (HUMAN).//
  - F-PLACE1008465//ZINC FINGER PROTEIN 31 (ZINC FINGER PROTEIN KOX29) (FRAGMENT).//0.00017:23: 43//HOMO SAPIENS (HUMAN).//P17040
- 30 F-PLACE1008488//HYPOTHETICAL PROTEIN UL61.//9.1e-05:204:30//HUMAN CYTOMEGALOVIRUS (STRAIN AD169).//P16818
  - F-PLACE1008524//HOMEOBOX PROTEIN HLX1 (HOMEOBOX PROTEIN HB24).//0.95:74:36//HOMO SAPIENS (HUMAN).//Q14774
  - F-PLACE1008531///// ALU SUBFAMILY SC WARNING ENTRY I/////3.1e-05:86:45//HOMO SAPIENS (HUMAN).//
    - F-PLACE1008532//HYPOTHETICAL 36.4 KD PROTEIN IN SMP1-MBA1 INTERGENIC REGION.//3.9e-21:62: 45//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38298
    - F-PLACE1008533//HYPOTHETICAL 86.2 KD PROTEIN C4G8.04 IN CHROMOSOME 1.//3.5e-06:118:29// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09830
- #0 F-PLACE1008568//NEURONATIN.//0.046;34;52//HOMO SAPIENS (HUMAN).//Q16517
  - F-PLACE1098584/HUNCHBACK PROTEIN (FRAGMENT)//0.94:30:43/LITHOBIUS FORFICATUS.//002030 F-PLACE1098603//NUCLEAR PORE COMPLEX PROTEIN NUP155 (NUCLEOPORIN NUP155) (155 KD NU-CLEOPORIN) (P140)//3.94-123:224:98//RATTUS NORVEGICUS (RATI.//P37199
- F-PLACE1008621//B-CELL GROWTH FACTOR PRECURSOR (BCGF-12 KD).//5.0e-05:31:67//HOMO SAPIENS (HUMAN).//P20931
- F-PLACE1008625//DISAGREGIN (PLATELET AGGREGATION ACTIVATION INHIBITOR).//0.87:17:52//ORNI-THODOROS MOUBATA (SOFT TICK).//P36235
  - F-PLACE1008626//METALLOTHIONEIN-I (MT-I).//0.77:33:36//SCYLLA SERRATA (MUD CRAB).//P02805
  - F-PLACE1008627//METALLOTHIONEIN-IIÌ (MT-III) (GROWTH INHIBITORY FACTOR) (GIF) (GIFB).//0.14:44: 31//HOMO SAPIENS (HUMAN).//P25713
- F-PLACE1008629

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- F-PLACE1008630//PROTAMINE Z3 (SCYLLIORHININE Z3).//0.78:33:36//SCYLIORHINUS CANICULA (SPOTTED DOGFISH) (SPOTTED CATSHARK).//P30258
- F-PLACE 1008643/INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN H4 PRECURSOR (ITI HEAVY CHAIN H4) (INTER-ALPHA-TRYPSIN INHIBITOR FAMILY HEAVY CHAIN-RELATED PROTEIN) (PLASMA KALLIKREIN SENSITIVE GLYCOPROTEIN 120) (PK-120)/I/7-8-30:220-47/IHOMO SAPIENS (HUMAN)//014624
  - F-PLACE1008650//PP1/PP2A PHOSPHATASES PLEIOTROPIC REGULATOR PRL1.//2.5e-10:106:31//ARABI-DOPSIS THALIANA (MOUSE-EAR CRESS).//Q42384

- F-PLACE1008693//BOWMAN-BIRK TYPE PROTEINASE INHIBITOR (MSTI).//1.0:36:38//MEDICAGO SCUTEL-LATA (SNAIL MEDIC).//P80321
- F-PLACE1008696//NADH-UBIQUINONE OXIDOREDUCTASE 23 KD SUBUNIT PRECURSOR (EC 1.6.5.3) (EC 1.6.99.3) (COMPLEX I-23KD) (CI-23KD) (TYKY SUBUNIT).//4.8e-14:47:80//HOMO SAPIENS (HUMAN).// 000217
- F-PLACE1008715//HYPOTHETICAL 13.4 KD PROTEIN IN ACT5-YCK1 INTERGENIC REGION. //0.66:105:24// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST). //P38834
  - F-PLACE1008748//HYPOTHETICAL 57.5 KD PROTEIN IN VMA7-RPS25A INTERGENIC REGION.//0.10:178: 26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53214
- 10 F-PLACE1008757//HYPOTHETICAL 10.1 KD PROTEIN IN RHSD-GCL INTERGENIC REGION (ORFD3).//0.60: 44:34//ESCHERICHIA COLI.//P33669
  - F-PLACE1008790//IMPORTIN ALPHA-6 SUBUNIT (KARYOPHERIN ALPHA-6 SUBUNIT) (IMPORTIN ALPHA S2).//3.0e-69:191:80//MUS MUSCULUS (MOUSE).//O35345 F-PLACE1008798//BACTERIOCIN LACTOBIN A. //1.0:34:41//LACTOBACILLUS AMYLOVORUS . //P80696
- F-PLACE1008807//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//0.91:77:36//HOMO SAPIENS (HU-15 MAN).//P08547
  - F-PLACE1008808//REC1 PROTEIN.//0.45:39:30//USTILAGO MAYDIS (SMUT FUNGUS).//P14746
    - F-PLACE1008813
  - F-PLACE1008851//VERY HYPOTHETICAL 11.8 KD PROTEIN IN KTR3-DUR1.2 INTERGENIC REGION.//1.0: 62:30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38309
  - F-PLACE1008854//HYPOTHETICAL 182.0 KD PROTEIN IN NMD5-HOM6 INTERGENIC REGION.//1.0:82:26// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47170
    - F-PLACE1008867//PATATIN T5 PRECURSOR (POTATO TUBER PROTEIN), //0.65:61:36//SOLANUM TUBERO-SUM (POTATO),//P15478
- 25 F-PLACE1008887//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.5e-56:180:54//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548 F-PLACE 1008902
  - F-PLACE1008920

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- F-PLACE1008925//HYPOTHETICAL 41.2 KD PROTEIN IN GAPA-RND INTERGENIC REGION.//0.90:77:33//ES-CHERICHIA COLI.//P76242
- F-PLACE1008934//HYPOTHETICAL PROTEIN IN ADHS 5'REGION (ORF3) (FRAGMENT), I/O. 14:77:45//GLU-CONOBACTER SUBOXYDANS //O05543
  - F-PLACE1008941//ZINC FINGER PROTEIN 141.//1.1e-17:45:95//HOMO SAPIENS (HUMAN).//Q15928
  - F-PLACE1008947//MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENP-B).//4.1e-14:136:39//MUS MUSCULUS (MOUSE).//P27790
  - F-PLACE1009020//ANNEXIN VII (SYNEXIN) (FRAGMENT).//0.74:37:48//BOS TAURUS (BOVINE).//P20072 F-PLACE1009027//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4L (EC 1.6.5.3).//0.39:57:36//BALAENOP-TERA MUSCULUS (BLUE WHALE).//P41301
- F-PLACE1009039
- F-PLACE1009045//HYPOTHETICAL 9.5 KD PROTEIN IN SPEA-METK INTERGENIC REGION (F83).//0.48:32: 40 43//ESCHERICHIA COLL//P46879 F-PLACE1009048

  - F-PLACE1009050
- F-PLACE1009060//HYPOTHETICAL 98.3 KD PROTEIN R10E12.1 IN CHROMOSOME III.//4.9e-23:244:31// 45 CAENORHARDITIS ELEGANS //P34552
  - F-PLACE1009090//50S RIBOSOMAL PROTEIN L35.//1.0:27:51//MYCOPLASMA GENITALIUM.//P47439 F-PLACE1009091
    - F-PLACE1009094//NEL-LIKE PROTEIN (FRAGMENT).//3.6e-15:180:30//HOMO SAPIENS (HUMAN).//Q92832 F-PLACE1009099//ZINC FINGER PROTEIN 27 (ZFP-27) (MKR4 PROTEIN) (FRAGMENT).//1.4e-94:228:71// MUS MUSCULUS (MOUSE).//P10077
  - F-PLACE1009110//HIRUDIN HV1 (BUFRUDIN).//1.0:49:34//HIRUDINARIA MANILLENSIS (BUFFALO LEECH).// PR1492
    - F-PLACE1009111//!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.4e-05:30:83//HOMO SAPIENS (HUMAN).//
- 55 F-PLACE1009113//ANNEXIN VII (SYNEXIN) (FRAGMENT).//0.032:40:52//BOS TAURUS (BOVINE).//P20072 F-PLACE1009130//HYPOTHETICAL PROTEIN KIAA0032.//3.3e-37:214:38//HOMO SAPIENS (HUMAN).// O15034
  - F-PLACE1009150//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.6e-32:56:76//HOMO SAPIENS (HUMAN).//

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- F-PLACE1009155/!!!! ALU SUBFAMILY SQ WARNING ENTRY!!!!//1.2e-17:101:57//HOMO SAPIENS (HU-MAN).//P39194
- F-PLACE1009158//HYPOTHETICAL PROTEIN HKRFX (J1I).//0.0058:73:42//HUMAN CYTOMEGALOVIRUS (STRAIN AD169).//P09711
- F-PLACE1009166//CYTOSOLIC PURINE 5'-NUCLEOTIDASE (EC 3.1.3.5),//0.0086:96:30//HOMO SAPIENS (HUMAN),/P49902
  - F-PLACE1009172//HYPOTHETICAL 8.7 KD PROTEIN IN GAPA-RND INTERGENIC REGION.//1.0:19:52//ES-CHERICHIA COLI.//978246
- F-PLACE1009174//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.1e-17:47:82//HOMO SAPIENS (HUMAN).// P39194

#### E-PLACE1009183

- F-PLACE1009186//HYPOTHETICAL 11.4 KD PROTEIN C13G6.04 IN CHROMOSOME I.//0.019:62:24// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09783
- F-PLACE1009190//PALMITOYL-COA HYDROLASE (EC 3.1.2.2) (LONG-CHAIN FATTY-ACYL-COA HYDROLA-SE) (FRAGMENT).//0.027:53:28//RATTUS NORVEGICUS (RAT).//P80250
  - F-PLACE1009200//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//5.4e-28:84:71//HOMO SAPIENS (HUMAN).// P39194
- F-PLACE1009230//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.8e-12:50:74//HOMO SAPIENS (HUMAN).//
  P39189
  - F-PLACE1009246//UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 7.8 KD PROTEIN (EC 1.10.2.2) (MITOCHONDRIAL HINGE PROTEIN) (CR7).//1.0:17:52//SOLANUM TUBEROSUM (POTATO).//P48504
    - F-PLACE1009298//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS35.//6.6e-41:177:53//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//P34110
- 25 F-PLACE1009308//KERATIN, ÜLTRA HIGH-SULFUR MATRIX PROTEIN (UHS KERATIN).//0.00034:108:33// HOMO SAPIENS (HUMAN).//P26371
  - F-PLACE1009319//PRESYNAPTIC DENSITY PROTEIN 95 (PSD-95),//5.3e-16:84:50//HOMO SAPIENS (HUMAN),//P78352
  - F-PLACE1009328//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//6.9e-82:263:67//HOMO SAPIENS (HU-MAN).//P08547
- F-PLACE1009335//60S RIBOSOMAL PROTEIN L32.//0.95:71:36//HOMO SAPIENS (HUMAN), MUS MUSCU-LUS (MOUSE), AND RATTUS NORVEGICUS (RAT).//P02433
  - F-PLACE1009338//TRANSCRIPTION FACTOR HES-5 (HAIRY AND ENHANCER OF SPLIT 5).//0.90:42:40// MUS MUSCULUS (MOUSE).//P70120
- F-PLACE1009988//BASIC PROLINE-RICH PEPTIDE IB-1//0.013:33:48//HOMO SAPIENS (HUMAN),//P04281 F-PLACE1009375//HYPOTHETICAL 88.1 KD PROTEIN K02D10.1 IN CHROMOSOME III.//0.0022:135:21// CAEMORHABDITIS ELEGANS.//P34492
  - F-PLACE1009388//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//4.8e-22:73:65//HOMO SAPIENS (HUMAN).// P39195
- 40 F-PLACE1009398//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2).//8.1e-83:223:65//HOMO SA-PIENS (HUMAN).//P51523
  - F-PLACE1009404//GLUTENIN, HIGH MOLECULAR WEIGHT SUBUNIT PW212 PRECURSOR.//0.047:145:29//
    TRITICUM AESTIVUM (WHEAT).//P08489
- F-PLACE1009410//TOXIN C13S1C1 PRECURSOR.//0.22:21:47//DENDROASPIS ANGUSTICEPS (EASTERN GREEN MAMBA).//P18329
  - F-PLACE1009434//NADH-UBIQUINONE OXIDOREDUCTASE SUBUNIT K (EC 1.6.5.3) (FRAGMENT).//0.81:61: 29//ANTHOCEROS FORMOSAE.//031791
    - F-PLACE1009443//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//9.1e-05:93:32//MUS MUS-CULUS (MOUSE).//Q62203
  - F-PLACE1009444/PHOSPHATIDYLINOSITOL 4-KINASE ALPHA (EC 2.7.1.67) (PI4-KINASE) (PTDINS-4-KI-NASE) (PI4-K-ALPHA)./I6.4e-15-41-97/IHOMO SAPIENS (HUMAN)./IP42356 F-PLACE1009459/IH/PDTHETICAL 423. KD PROTEIN C12G[12 11] CIN CHROMOSOME I.//0.0011:119:31//
    - SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09874

      F-PLACE1009468//PHOSPHOLIPASE A-2-ACTIVATING PROTEIN (PLAP).//4.2e-34:101:75//RATTUS NOR-
- 59 VEGICUS (RAT),//P54319 F-PLACE109476/IDNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS),//0.086:21:52//HO-MO SAPIENS (HUMAN),//P30808
  - F-PLACE1009477

- F-PLACE1009493//HYPOTHETICAL 127.3 KD PROTEIN B0416.1 IN CHROMOSOME X.//1.4e-18:138:39// CAENORHABDITIS ELEGANS //O11069
- F-PLACE1009524//ARF NUCLEOTIDE-BINDING SITE OPENER (ARNO PROTEIN) (ARF EXCHANGE FACTOR),//9.4e-80:155:85//HOMO SAPIENS (HUMAN),//Q99418
- 5 F-PLACE1009539//GTP-BINDING NUCLEAR PROTEIN RAN/TC4.//1.0:76:26//GIARDIA LAMBLIA (GIARDIA IN-TESTINALIS).//P38543
  - F-PLACE1009542//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.00016:31:77//HOMO SAPIENS (HUMAN).//
- F-PLACE1009571//ATP SYNTHASE B CHAIN (EC 3.6.1.34) (SUBUNIT I).//0.88:116:29//STREPTOCOCCUS PNEUMONIAE.//059952
  - F-PLACE1009581//50S RIBOSOMAL PROTEIN L32.//0.00023:37:51//RHODOBACTER CAPSULATUS (RHODOPSEUDOMONAS CAPSULATA).//P30788
- F-PLACE1009595
  F-PLACE1009596//HYPOTHETICAL 40.4 KD TRP-ASP REPEATS CONTAINING PROTEIN C14B1.4 IN CHRO-
- F-PLACE1009596//HYPOTHETICAL 40.4 KD TRP-ASP REPEATS CONTAINING PROTEIN C14B1.4 IN CHRO-MOSOME III.//2.1e-36:116:49//CAENORHABDITIS ELEGANS.//Q17963
  - F-PLACE1009607//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.8e-43:73:69//HOMO SAPIENS (HUMAN).// P39188
    - F-PLACE1009613

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F-PLACE1009886

- F-PLACE1009621//TRANSCRIPTION FACTOR BTF3 HOMOLOG 2.//0.91:29:44//HOMO SAPIENS (HUMAN).//
- F-PLACE1009622//MATERNAL EFFECT PROTEIN STAUFEN.//1.3e-22:132:47//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P25159
  - F-PLACE1009637//HYPOTHETICAL 18.1 KD PROTEIN IN CFXA 3'REGION.//0.30:28:57//BACTEROIDES VUL-GATUS.//P30905
- 25 F-PLACE1009639//LIPASE MODULATOR PRECURSOR (LIPASE HELPER PROTEIN).//0.23:79:31//PSEU-DOMONAS AERUGINOSA.//Q04591
  - F-PLACE1099659/MEMRANE-ASSOCIATED PROTEIN HEM-2 (BRAIN PROTEIN H19) (MH19) (FRAG-MENT), //3.9e-126-227:96/MUS MUSCULUS (MOUSE), //P28660 F-PLACE1099865//G KAPPA CHAIN V-I REGION (HAU), //5.52:89-35/H/DMO SAPIENS (HUMAN), //P01600
- 59 F-PLACE1009670/CYCLOMALTODEXTRIN GLUCANOTRANSFERASE PRECURSOR (EC 2.4.1.19) (CYCLO-DEXTRIN-GLYCOSYLTRANSFERASE) (CGTASE).//0.16:114:29//PAENIBACILLUS MACERANS (BACILLUS MACERANS).//B13856
  - F-PLACE1009708//HYPOTHETICAL 143.3 KD TRP-ASP REPEATS CONTAINING PROTEIN C12G12.13C IN CHROMOSOME I.//9.6e-19:156:36//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09876
- 35 F-PLACE1009721//MSF1 PROTEIN.//7.7e-23:176:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).// P35200
  - F-PLACE1009731//AIG1 PROTEIN.//1.1e-09:91:43//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).// P54120
- F-PLACE1009763//HYPOTHETICAL 48.9 KD PROTEIN C24H6.12C IN CHROMOSOME I.//8.3e-42:171:51//
  SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09765
  - F-PLACE1009794//C-HORDEIN (CLONE PC HOR1-3) (FRAGMENT).//0.99:36:33//HORDEUM VULGARE (BAR-LEY).//P17991
  - F-PLACE1009798//HYPOTHETICAL PROTEIN C22F3.14C IN CHROMOSOME I (FRAGMENT).//2.6e-34:191: 38//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09779
- 45 F-PLACE1009845/WEB1 PROTEIN (PROTEIN TRANSPORT PROTEIN SEC31).//2.26-19:190:33//SACCHA-ROMYCES CEREVISIAE (BAKER'S YEAST).//P38968 F-PLACE1009861/JCATHEPSIN B PRECURSOR (EC 3.4.22.1).//4.46-20:171:33//BOS TAURUS (BOVINE).//
  - P07688
  - F-PLACE1009879//HYPOTHETICAL 8.7 KD PROTEIN IN RPL22-RPL23 INTERGENIC REGION (ORF70).//0.99: 30:33//ASTASIA LONGA (EUGLENOPHYCEAN ALGA).//P34779
    - F-PLACE1009888//NONSTRUCTURAL POLYPROTEIN [CONTAINS: NONSTRUCTURAL PROTEIN NSP4] (FRAGMENT),//1.0:33:42//WESTERN EQUINE ENCEPHALITIS VIRUS.//P13896
- F.-PLACE 1009908//HYPOTHETICAL GTP-BINDING PROTEIN C3F10.16C IN CHROMOSOME 1.//3.1e-42:205: 55 46//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10190 F.-PI ACE 1009921
  - F-PLACE1009924//ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6).//0.70:128:29//TRYPANOSOMA BRU-CEI BRUCEI./P24499

- F-PLACE1009925//ATP SYNTHASE D CHAIN, MITOCHONDRIAL (EC 3.6.1.34).//0.99:111:27//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P30902
- F-PLACE1009935//HYPOTHETICAL PROTEIN MJ0258.//0.063:75:32//METHANOCOCCUS JANNASCHII.//
- F-PLACE1009947//NEUROGRANIN (NG) (P17) (B-50 IMMUNOREACTIVE C-KINASE SUBSTRATE) (BICKS) (FRAGMENT).//0.33:51:45/BOS TAURUS (BOVINE).//P35722
  - F-PLACE1009971//MIPP PROTEIN (MURINE IAP-PROMOTED PLACENTA-EXPRESSED PROTEIN).//0.022:
  - 84:27//MUS MUSCULUS (MOUSE).//P28575 F-PLACE1009992//BONE MORPHOGENETIC PROTEIN 1 PRECURSOR (EC 3.4.24.-) (BMP-1).//0.00011:35:
- 51//HOMO SAPIENS (HUMAN).//P13497
  F-PLACE1009995//TROPOMYOSIN, SMOOTH MUSCLE/FIBROBLAST CTM1.//0.052:185:22//CIONA INTESTI-
  - NALIS //007068 F-PLACE1009997/TRANSCRIPTION ELONGATION FACTOR S-II (RNA POLYMERASE II ELONGATION FAC-TOR DMS-II) (TFIIS) //0.68:98:28//DROSOPHILA MELANOGASTER (FRUIT FLY).//P20232
- 15 F-PLACE1010023/HVPOTHETICAL 83.8 KD PROTEIN C27F2.7 IN CHROMOSOME III.//6.6e-06:111:32//
- CAENORHABDITIS ELEGANS.//018262

  F-PLACE1010031//HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION.//0.0024:72:33//
  AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMIPPV).//P41479
  - F-PLACE1010053//HYPOTHETICAL PROTEIN HI0593.//0.83:24:45//HAEMOPHILUS INFLUENZAE.//P44022 F-PLACE1010069
- F-PLACE1010069 F-PLACE1010074/NACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS5.//0.00027:192:28//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST) //.092331
  - F-PLACE1010076//HUNCHBACK PROTEIN (FRAGMENT).//0.80:39:30//SCIARA COPROPHILA (FUNGUS GNAT).//001790
- 25 F-PLACE1010083//RHO-GAP HEMATOPOIETIC PROTEIN C1 (P115) (KIAA0131).//2.7e-48:177:46//HOMO SA-PIENS (HUMAN).//P98171
  - F-PLACE1010088//UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 11 (EC 3.1.2.15) (UBIQUITIN THI-OLESTERASE 11) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 13) (DEUBIQUITINATING ENZYME 11) (KIAAQ055)://7.96-07.55.43//HOMO SAPIENS (HUMAN).//P40818
- F-PLACE1010096/I/100 KD PROTEIN (EC 6.3.2-)//1.0e-107:232:90//RATTUS NORVEGICUS (RAT).//G62671 F-PLACE1010102//DNA-DIRECTED RNA POLYMERASE SUBUNIT N (EC 2.7.7.6).//1.0.33:45//METHANOCOC-CUS JANNASCHII.//057649
  - F-PLACE1010105//RING CANAL PROTEIN (KELCH PROTEIN).//1.2e-47:200:46//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//004652
- 35 F-PLACE1010106//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS: REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE].//1.2e-14:94:41//MUS MUSCULUS (MOUSE).//P11369
  - F-PLACE1010134//HYPOTHETICAL 171.5 KD HELICASE IN NUT1-ARO2 INTERGENIC REGION.//4.0e-28:78: 76//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53115
- F-PLACE1010148//GAR2 PROTEIN.//2.6e-05:180:26//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//P41891
  - F-PLACE1010152/UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 64E (EC 3.1.2.15) (UBIQUITIN THI-OLESTERASE 64E) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 64E) (DEUBIQUITINATING ENZYME 64E)/I/2 1e-59:227-54/DROSOPHILA MELANOGASTER (FRUIT FLY)//024574
- F-PLACE1010181/MALE SPECIFIC SPERM PROTEIN MST87F.//0.39:12:58//DROSOPHILA MELANOGASTER

  (FRUIT FLY).//P08175
- F-PLACE1010194/ISPLICING FACTOR, ARGININE/SERINE-RICH 2 (SPLICING FACTOR SC35) (SC-35) (SPLICING COMPONENT, 35 KD) (PR264 PROTEIN),///14e-07-95-43//CALLUS GALLUS (CHICKEN),///P30352 F-PLACE1010202//TRISTETRAPROLINE (TTP) (TIS11A) (TIS11) (ZFP-36),///.0.094:109:29//RATTUS NORVEGICUS (RAT),///P47973
  - F-PLACE 1010231/ILANTIBIOTIC NISIN A PRECURSOR //0.99:42:35//LACTOCOCCUS LACTIS (SUBSP. LACTIS) (STREPTOCOCCUS LACTIS) //P13068
    - F-PLACE1010261//SEGREGATION DISTORTER PROTEIN.//6.0e-71:201:62//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//P25722
- F-PLACE1010270

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- 55 F-PLACE1010274//HYPOTHETICAL 16.2 KD PROTEIN C4F8.01 IN CHROMOSOME I.//4.4e-08:100:26// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O14177
  - F-PLACE1010293//IIII ALU SUBFAMILY J WARNING ENTRY IIII//3.9e-26:94:64//HOMO SAPIENS (HUMAN).// P39188

- F-PLACE1010310//SYNAPSINS IA AND IB //5.7e-09:89:37//RATTUS NORVEGICUS (RAT) //P09951
- F-PLACE1010321//IMMEDIATE-EARLY PROTEIN IE180.//0.033:145:31//PSEUDORABIES VIRUS (STRAIN KA-PLAN) (PRV).//P33479
- F-PLACE1010324//MAST CELL DEGRANULATING PEPTIDE (MCDP) (MCD).//0.60:25:48//MEGABOMBUS PENNSYLVANICUS (AMERICAN COMMON BUMBLEBEE).//P04567
- F-PLACE1010329//TOXIN S5C10.//1.0:39:33//DENDROASPIS JAMESONI KAIMOSAE (EASTERN JAMESON'S MAMBA) //P01419
  - F-PLACE1010341//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//0.0049:49:55//HOMO SAPIENS (HUMAN).//
- F-PLACE1010362//NARIANT-SURRACE-GLYCOPROTEIN PHOSPHOLIPASE C (EC 3.1.4.47) (VSG LIPASE) (GLYCOSYLPHOSPHATIDYLINOSITOL-SPECIFIC PHOSPHOLIPASE C) (GPL-PLC).//0.0034:89:30// TRYPANOSOMA CRUZI.//015886
  - F-PLACE1010364//NADH-UBIQUINONE OXIDOREDUCTASE B17 SUBUNIT (EC 1.6.5.3) (EC 1.6.99.3) (COM-PLEX I-B17) (CI-B17).//1.0:40:35//SUS SCROFA (PIG).//Q29259
- 15 F-PLACE1010383

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- F-PLACE1010401//140 KD NUCLEOLAR PHOSPHOPROTEIN (NOPP140).//0.10:174:22//RATTUS NORVEGI-CUS (RAT) //P41777
  - F-PLACE1010481/HYPOTHETICAL 71.9 KD PROTEIN B0285.5 IN CHROMOSOME III.//1.5e-21:170:35// CAENORHABDITIS ELEGANS.//P46555
- F-PLACE1010491//HYPOTHETICAL 13.5 KD PROTEIN IN MOB1-SGA1 INTERGENIC REGION.//1.0:31:41// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40490
  - F-PLACE1010492//HYPOTHETICAL 42.3 KD PROTEIN C12G12.11C IN CHROMOSOME I.//0.77:97:30// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q09874
  - F-PLACE1010522//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.74:45:37//HOMO SAPIENS
    - F-PLACE 1010529//DELTA 1-PYRROLINE-5-CARBOXYLATE SYNTHETASE (P5CS) [CONTAINS: GLUTAMATE 5-KINASE (EC 2.7.2.11) (GAMMA-GLUTAMYL KINASE) (GK); GAMMA-GLUTAMYL PHOSPHATE REDUCTASE (GPR) (EC 1.2.1.41) (GLUTAMATE-5-SEMIALDEHYDE DEHYDROGENASE) (GLUTAMYL-GAMMA-SEMIALDEHYDE DEHYDROGENASE) (FLUTAMYL-GAMMA-SEMIALDEHYDE DEHYDROGENASE) (FLUTAMYL-GAMMA-SEMIALDEHYDROGENASE) (FLUTAMYL-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-GAMMA-G
- 30 F-PLACE1010547//HYPOTHETICAL 31.0 KD PROTEIN IN BUD9-RME1 INTERGENIC REGION.//0.17:68:39// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53227
  - F-PLACE1010562//CHLOROPLAST 50S RIBOSOMAL PROTEIN L33.//0.50:48:29//PORPHYRA PURPUREA.// P51255
- F-PLACE1010579/IHYPOTHETICAL PROTEIN HI1571.//0.29:37-43/IHAEMOPHILUS INFLUENZAE./IP44260
  F-PLACE1010580/IPUTATIVE ATP-DEPENDENT RINA HELICASE C12C2.06.I/3.3e-38:178:48/ISCHIZOSAC-CHAROMYCES POMBE (FISSION YEAST).//009747
  - F-PLACE1010599//PEROXISOMAL MEMBRANE PROTEIN PER10 (PEROXIN-14).//4.6e-17:192:31//PICHIA ANGUSTA (YEAST) (HANSENULA POLYMORPHA).//P78723
- F-PLACE1010616//HYPOTHETICAL 9.2 KD PROTEIN IN RNPA 3'REGION.//0.44:32:37//PSEUDOMONAS PUT-40 IDA.//P25753
  - F-PLACE1010622//A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR.//5.0e-06:102:42//SACCHAROMY-CES CEREVISIAE (BAKER'S YEAST).//P32323
  - F-PLACE1010624//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.00036:134:321/HOMO SAPIENS (HUMAN).//P10162
- 45 F-PLACE1010628
  - F-PLACE1010629//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.7e-12:37:81//HOMO SAPIENS (HUMAN).// P39194
    - F-PLACE1010630
  - F-PLACE1010631//WNT-5B PROTEIN (FRAGMENT).//0.49:62:30//EUMECES SKILTONIANUS (WESTERN SKINK) //P28118
  - F-PLACE1010661//MATERNAL EXUPERANTIA 2 PROTEIN.//1.0:95:30//DROSOPHILA PSEUDOOBSCURA (FRUIT FLY).//Q24617
    - F-PLACE1010662/JUDP-GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (EC 2.4.1.-)
      (DUGT).//3.2e-05:117:24//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q09332
- 55 F-PLACE1010702//ZINC FINGER PROTEIN 195.//1.4e-62:117:62//HOMO SAPIENS (HUMAN).//O14628 F-PLACE1010714
  - F-PLACE1010720//CHROMOSOME ASSEMBLY PROTEIN XCAP-C.//1.1e-64:176:76//XENOPUS LAEVIS (AF-RICAN CLAWED FROG).//P50532

- F-PLACE1010739//TAT PROTEIN (TRANSACTIVATING REGULATORY PROTEIN) (FRAGMENT).//0.97:31:41// HUMAN IMMUNODEFICIENCY VIRUS TYPE 1 (BH5 ISOLATE) (HIV-1).//P04612
- F-PLACE1010743//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//3.8e-05:253:30//MUS MUSCULUS (MOUSE).//P05143
- F-PLACE1010761//HYPOTHETICAL 37.0 KD PROTEIN B0495.8 IN CHROMOSOME II.//1.5e-14:175:25// CAENORHABDITIS ELEGANS.//Q09217
  - F-PLACE1010771//TRANSCRIPTIONAL REGULATOR PROTEIN HCNGP.//1.3e-120:216:89//MUS MUSCULUS (MOUSE).//Q02614
- F-PLACE1010786//CENTROSOMIN (ARROW PROTEIN).//0.97:133:24//DROSOPHILA MELANOGASTER
  (FRUIT FLY).//P54623
  - F-PLACE1010800//HYPOTHETICAL 31.7 KD PROTEIN IN TRAX-FINO INTERGENIC REGION (ORFC).// 0.0060:111:31//ESCHERICHIA COLL.//Q99390
  - F-PLACE1010802//UREASE ACCESSORY PROTEIN UREI.//0.82:44:29//BACILLUS SP. (STRAIN TB-90).// Q07415
- 15 F-PLACE1010811//CYTOCHROME C-551 (C551).//0.99:42:38//ECTOTHIORHODOSPIRA HALOCHLORIS.// P38587
  - F-PLACE1010833//CALTRACTIN, ISOFORM 1 (CENTRIN).//2.8e-09:90:34//HOMO SAPIENS (HUMAN).// P41208
  - F-PLACE1010856//MOLT-INHIBITING HORMONE (MIH).//1.0:32:37//PROCAMBARUS CLARKII (RED SWAMP CRAYFISH).//P55848
    - F-PLACE1010857//IG ALPHA-1 CHAIN C REGION.//0.49:73:34//GORILLA GORILLA GORILLA (LOWLAND GORILLA).//P20758
    - F-PLACE1010870//ZINC FINGER PROTEIN 91 (ZINC FINGER PROTEIN HTF10) (HPF7).//1.2e-56:173:58//HO-MO SAPIENS (HUMAN).//Q05481
- 25 F-PLACE1010877//HEAT SHOCK PROTEIN 82.//0.13:130:25//ZEA MAYS (MAIZE).//Q08277
  - F-PLACE1010891//HYPOTHETICAL 8.2 KD PROTEIN IN BLTR-SPOIIIC INTERGENIC REGION.//0.95:51:27// BACILLUS SUBTILIS.//P54436
  - F-PLACE1010896//SERINE/THREONINE-PROTEIN KINASE PTK1/STK1 (EC 2.7.1.).//0.98:71:30//SACCHA-ROMYCES CEREVISIAE (BAKER'S YEAST).//P36002
- F-PLACE1010900/HYPOTHETICAL PROTEIN HI0840.//1.0:42:30//HAEMOPHILUS INFLUENZAE.//P44897 F-PLACE1010916//KERATIN, HIGH-SULFUR MATRIX PROTEIN, IIIB3.//0.060:59:35//OVIS ARIES (SHEEP).// P02444
  - F-PLACE1010917//E2 GLYCOPROTEIN PRECURSOR (SPIKE GLYCOPROTEIN) (PEPLOMER PROTEIN).//
    0.71:141:24//BOVINE CORONAVIRUS (STRAIN L9).//P25191
- 35 F-PLACE1010925//HYPOTHETICAL 8.1 KD PROTEIN.//1.0:17:58//THERMOPROTEUS TENAX VIRUS 1 (STRAIN KRA1) (TTV1).//P19285
  - F-PLACE1010926//HYPOTHETICAL PROLINE-RICH PROTEIN KIAA0269.//0.011:51:45//HOMO SAPIENS (HU-MAN) //092558
- F-PLACE1010942//EPIDERMAL GROWTH FACTOR RECEPTOR SUBSTRATE SUBSTRATE 15 (PROTEIN EPS15)://3.1e-09:64:37//MUS MUSCULUS (MOUSE)://P42567
  - F-PLACE:1010944//GAP JUNCTION ALPHA-3 PROTEIN (CONNEXIN 44) (CX44).//0.17:71:38//BOS TAURUS (BOVINE).//P41987
    F-PLACE:1010947
- F-PLACE1010954//TROPOMYOSIN ALPHA CHAIN, SKELETAL MUSCLE.//0.011:144:26//HOMO SAPIENS
- F-PLACE1010960//ACTIN-LIKE PROTEIN 13E.//1.1 e-60:136:52//DROSOPHILA MELANOGASTER (FRUIT FLY) //P45890
  - F-PLACE1010965

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- F-PLACE1011026//PERIOD CLOCK PROTEIN (FRAGMENT).//1.0:64:31//DROSOPHILA ANANASSAE (FRUIT FLY) //003293
- F-PLACE1011032//RIBONUCLEASE HI (EC 3.1.26.4) (RNASE HI) (RIBONUCLEASE H) (RNASE H).//1.0:32:37// SALMONELLA TYPHIMURIUM.//P23329
  - F-PLACE1011041//HOMEOBOX PROTEIN VAB-7,//0.36:65:30//CAENORHABDITIS ELEGANS.//Q93899
- F-PLACE1011048/I-P-HOSPHATIDYLINOSITOL-4,5-BISPHOSPHATE PHOSPHODIESTERASE BETA 1 (EC 3.1.4.11) (PLC-BETA-1) (PHOSPHOLIPASE C-BETA-1) (PLC-1) (PLC-154).//1.3e-22:58:93//RATTUS NORVEGICUS (RAT)./P10887
  - F-PLACE1011054//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//1.6e-07:38:73//HOMO SAPIENS (HUMAN).// P39195

- F-PLACE1011056//HISTONE H1.//2.2e-10:109:41//PISUM SATIVUM (GARDEN PEA).//P08283
- F-PLACE1011057
  F-PLACE1011090/HYPOTHETICAL 33.8 KD PROTEIN IN TWT1-FLO5 INTERGENIC REGION.//1.8e-07:133:
- 32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38892 F-PLACE1011109//ELONGATION FACTOR G. MITOCHONDRIAL PRECURSOR (MEF-G).//5.4e-25:63:88//RAT-
- TUS NORVEGICUS (RAT).//Q07803

  F.PI ACF1011114//PUTATIVE\_ATP-DEPENDENT\_RNA\_HELICASE\_C1F7.02C.//8.4e-31:157:45//SCHIZOSAC-
  - CHAROMYCES POMBE (FISSION YEAST).//Q09916
- F-PLACE1011133//SERUM AMYLOID P-COMPONENT PRECURSOR (SAP) (9.5S ALPHA-1-GLYCOPRO-TEIN)//0.92:58:31//HOMO SAPIENS (HUMAN)//P02743
  - F-PLACE1011143/IPROBABLE E5 PROTEIN./I0.24-42:35/I/HUMAN PAPILLOMAVIRUS TYPE31./IP17385 F-PLACE1011160/IEARLY NODULIN 55-2 PRECURSOR (N-55-2) (NODULIN-315)./I0.88:98:27/I/GLYCINE MAX (SOYBEAN)./IO02917
- F-PLACE1011165//HISTIDINE-RICH PROTEIN.//0.013:13:76//PLASMODIUM FALCIPARUM (ISOLATE FCM17 / SENEGAL).//P14586
- F-PLACE1011185//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.4e-13:98:50//HOMO SAPIENS (HUMAN).// P39188
  - F-PLACE1011203
  - F-PLACE1011214//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:48:27//EQUUS ASINUS (DONKEY).//
  - F-PLACE1011219//PROBABLE OXIDOREDUCTASE (EC 1.-..-).//1.9e-15:162:31//STREPTOMYCES ANTIBI-OTICUS.//003326
    - F-PLACE1011221//ANTITHROMBIN-III HOMOLOG.//0.84:74:33//FOWLPOX VIRUS (ISOLATE HP-438[MU-NICHI).//P14369
- 25 F-PLÄCE1011229//UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 4 (EC 3.1.2.15) (UBIQUITIN THIOLESTE-RASE 4) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 4) (DEUBIQUITINATING ENZYME 4) (UBIQUI-TOUS NUCLEAR PROTEIN HOMOLOG)//3.56-86:218:68//HOMO SAPIENS (HUMAN)//013107
  - F-PLACE1011263//ANKYRIN, BRAIN VARIANT 2 (ANKYRIN B) (ANKYRIN, NONERYTHROID) (FRAGMENT).// 3.0e-07:99:36//HOMO SAPIENS (HUMAN).//Q01485
- 30 F-PLACE1011273

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- F-PLACE1011291//PROTEIN KINASE C SUBSTRATE 80 KD PROTEIN (FRAGMENTS).//0.011:36:50//RATTUS NORVEGICUS (RAT).//P20468
  - F-PLACE1011296//HOMEOBOX PROTEIN DLX-6.//0.76:55:32//BRACHYDANIO RERIO (ZEBRAFISH) (ZEBRA DANIO).//Q98877
- 35 F-PLACE1011310//ATP SYNTHASE PROTEIN 9, MITOCHONDRIAL (EC 3.6.1.34) (LIPID-BINDING PROTEIN).// 0.46:43:44//PETUNIA SP. (PETUNIA).//Q07060
  - F-PLACE1011325//HYPOTHETICAL 222.8 KD PROTEIN C1F3.06C IN CHROMOSOME I.//0.00021:171:27// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10411
  - F-PLACE1011332//DNA-DAMAGE-REPAIR/TOLERATION PROTEIN DRT101 PRECURSOR.//7.3e-27:113:52// ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).//Q05211
    - F-PLACE1011340//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.6e-07:40:62//HOMO SAPIENS (HUMAN).//
  - F-PLACE1011371//INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN H2 PRECURSOR (ITI HEAVY CHAIN H2) //2 2e-54:227:44//MUS MUSCULUS (MOUSE) //061703
- 45 F-PLACE1011375//PROBABLE E5 PROTEIN.//0.93:28:57//HUMAN PAPILLOMAVIRUS TYPE 51.//P26553
- F-PLACE1011399//HISTONE H2B-IV.//0.19:129:27//VOLVOX CARTERI.//P16868 F-PLACE1011419
  - F-PLACE1011419
    F-PLACE1011433//ZINC FINGER PROTEIN GLI3 (FRAGMENT).//3.4e-05:133:24//GALLUS GALLUS (CHICK-EN).//P55879
- 50 F-PLACE1011452//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//3.9e-25:76:63//HOMO SAPIENS (HU-MAN)./P08547
  - F-PLACE1011465//ECTODERMAL DYSPLASIA PROTEIN (EDA PROTEIN).//0.97:36:41//HOMO SAPIENS (HU-MAN).//092838
- F-PLACE1011472//METALLOTHIONEIN-1 (CUMT-1).//0.084:55:30//HOMARUS AMERICANUS (AMERICAN LOBSTER).//P29499
  - F-PLACE 1011477//CELL SURFACE GLYCOPROTEIN 1 PRECURSOR (OUTER LAYER PROTEIN B) (S-LAYER PROTEIN 1).//0.028:129:34//CLOSTRIDIUM THERMOCELLUM.//Q06852
  - F-PLACE1011492//NON-GREEN PLASTID TRIOSE PHOSPHATE TRANSLOCATOR PRECURSOR (CTPT).//

2.9e-13:147:31//BRASSICA OLERACEA (CAULIFLOWER).//P52178

F-PLACE1011503//PUTATIVE FERREDOXIN-LIKE PROTEIN IN PURL-DPJ INTERGENIC REGION (086).//0.66: 32:40//ESCHERICHIA COLL//PE2102

F-PLACE1011520

F-PLACE1011563//LORICRIN.//0.00023:112:39//HOMO SAPIENS (HUMAN).//P23490

F-PLACE1011567//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//9.2e-31:78:76//HOMO SAPIENS (HUMAN).// P39195

F-PLACE1011576//ZINC FINGER PROTEIN 91 (ZINC FINGER PROTEIN HTF10) (HPF7).//1.5e-32:45:86//HO-MO SAPIENS (HUMAN).//Q05481

F-PLACE1011586/IN-TYPE CALCIUM CHANNEL ALPHA-18 SUBUNIT (OMEGA-CONOTOXIN-SENSITIVE N TYPE, BRAIN CALCIUM CHANNEL ALPHA-1 SUBUNIT)/IO:26:8137/iH/OMO SAPIENS (HUMAN)/IO00975 F-PLACE1011635//IMMEDIATE-EARLY PROTEIN IE180./IO:00045:170:30/PSEUDORABIES VIRUS (STRAIN INDIAMA-FUNKHAUSER REFCKER IPRVI)/IP11675.

INDIANA-FUNKHAUSER /BECKER) (PRV).//P11675

15 F-PLACE1011643//CUTICLE COLLAGEN 40.//1.0:128:32//CAENORHABDITIS ELEGANS.//P34804

F-PLACE1011646//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-15:44:63//HOMO SAPIENS (HUMAN).//

F-PLACE1011649//HYPOTHETICAL PROTEIN F-215.//0.48:106:34//HUMAN ADENOVIRUS TYPE 2.//P03291 F-PLACE1011650

F-PLACE1011664//CROOKED NECK PROTEIN.//1.2e-79:201:68//DROSOPHILA MELANOGASTER (FRUIT FLY).//P17886

F-PLACE1011675//FERREDOXIN.//1.0:44:29//METHANOCOCCUS THERMOLITHOTROPHICUS.//P21305 F-PLACE1011682//HYPOTHETICAL 7.0 KD PROTEIN IN PRS26A-COX4 INTERGENIC REGION.//1.0:40:22// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53088

F-PLACE1011719//NEUROTOXIN TX2-6.//0.90:31:35//PHONEUTRIA NIGRIVENTER (BRAZILIAN ARMED SPI-DER).//P29425

F.-PLACE1011725/INUCLEOBINDIN PRECURSOR (NUCB1) (BONE 63 KD CALCIUM-BINDING PROTEIN).//
0.0065:125:25/IRATTUS NORVEGICUS (RAT).//Q63083
F.-PLACE1011729//SRY-RELATED PROTEIN LG27 (FRAGMENT).//0.97-48:39//EUBLEPHARIS MACULAR-

F-PLACE1011/29//SRY-RELATED PROTEIN LG2/ (FRAGMENT)://0.97/48/39//EUBLEPHARIS MACULA
30 IUS //P40654

F-PLACE1011749

F-PLACE1011762//D-BINDING PROTEIN (DBP) (ALBUMIN D BOX-BINDING PROTEIN).//0.028:91:39//MUS MUSCULUS (MOUSE).//Q60925 F-PLACE1011778

F-PLACE1011783//EMBRYONIC GROWTH/DIFFERENTIATION FACTOR 1 PRECURSOR (GDF-1).//0.97:48: 43//MUS MUSCULUS (MOUSE).//P20863

F-PLACE1011858//COLLAGEN 1(X) CHAIN PRECURSOR./I/0.0027:154:33//BOS TAURUS (BOVINE).//P23206 F-PLACE1011874//BACTERIOCHLOROPHYLL A PROTEIN (BCHL A PROTEIN) (BCP).//1.0:60.26//PROSTHE-COCHL ORLS AFSTIJABII //P11741

40 F-PLACE1011875//HYPOTHETICAL 6.6 KD PROTEIN IN GP54-ALT INTERGENIC REGION.//0.99:34:35//AC-TERIOPHAGE T4 //P39495

F-PLACE1011891//SMOOTHELIN.//0.018:122:31//HOMO SAPIENS (HUMAN).//P53814

F-PLACE1011896//SKIN SECRETORY PROTEIN XP2 PRECURSOR (APEG PROTEIN).//6.3e-09:203:35// XENOPUS LAEVIS (AFRICAN CLAWED FROG).//P17437

45 F-PLACE1011922//CRYPTDIN-RELATED PROTEIN 4C-2 PRECURSOR (CRS4C).//0.067:37:48//MUS MUSCU-LUS (MOUSE).//P50715

F-PLACE1011923//SERINE/THREONINE-PROTEIN KINASE SNK (EC 2.7.1.-) (SERUM INDUCIBLE KINASE).// 1.5e-83:175:89//MUS MUSCULUS (MOUSE).//P53351

F-PLACE1011962//MATING-TYPE PHEROMONE BAP1(2) PRECURSOR.//0.50:46:41//SCHIZOPHYLLUM COMMUNE (BRACKET FUNGUS).//Q02593

F-PLACE1011964//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG.//1.6e-05:47:51//NYCTICEBUS COU-CANG (SLOW LORIS).//P08548

F-PLACE1011982//APICAL MEMBRANE ANTIGEN 1 PRECURSOR (MEROZOITE SURFACE ANTIGEN).//0.98: 83:31//PLASMODIUM FRAGILE //P22622

55 F-PLACE1011995

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F-PLACE1012031/HYPOTHETICAL PROTEIN KIAA0254.//0.032:62:33/HOMO SAPIENS (HUMAN).//Q92543 F-PLACE2000003///III ALU SUBFAMILY SP WARNING ENTRY !!!!//5.4e-18:63:73/HOMO SAPIENS (HUMAN).// P39193

- F-PLACE2000008/JANNEXIN VII (SYNEXIN) (FRAGMENT).//0.14:20:50/JBOS TAURUS (BOVINE).//P20072
  F-PLACE2000007/PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.0045:176:30/JMUS MUSCULUS (MOUSE).//P05143
- F-PLACE2000011//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.6e-25:57:78//HOMO SAPIENS (HUMAN).// P39194
  - F-PLACE2000014//HYPOTHETICAL HELICASE C28H8.3 IN CHROMOSOME III.//0.00013:237:27// CAENORHARDITIS ELEGANS.//009475
  - F-PLACE2000015//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.4e-33:60:80//HOMO SAPIENS (HUMAN).// P39193
- 10 F-PLACE2000017/I/FOLATE RECEPTOR BETA PRECURSOR (FR-BETA) (FOLATE RECEPTOR 2) (FOLATE RECEPTOR, FETAL/PLACENTAL) (PLACENTAL FOLATE-BINDING PROTEIN) (FBP).//10:83:31/I/HOMO SAPI-ENS (HUMAN)./P14207
  - F-PLACE2000021//EPHRIN TYPE-A RECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN KI-NASE RECEPTOR CEK8).//0.99:103:26//GALLUS GALLUS (CHICKEN).//Q07496
- 15 F-PLACE2000030//MALE SPECIFIC SPERM PROTEIN MST84DA.//0.69:29:44//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q01642
  - F-PLACE2000033//PROBABLE OXIDOREDUCTASE (EC 1.-.-.).//1.1e-05:74:41//STREPTOMYCES ANTIBIOTI-
  - F-PLACE2000034//AXONIN-1 PRECURSOR (AXONAL GLYCOPROTEIN TAG-1) (TRANSIENT AXONAL GLYC-OPROTEIN 1).//6.7e-18:191:35//HOMO SAPIENS (HUMAN).//Q02246
  - F-PLACE2000039//DYNEIN HEAVY CHAIN, CYTOSOLIC (DYHC) (MAP 1C).//4.7e-80:163:96//RATTUS NOR-VEGICUS (RAT).//P38650
    - F-PLACE2000047///!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!//6.4e-06:63:49//HOMO SAPIENS (HU-MAN).//P39191
- 25 F-PLACE2000050//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//3.2e-22:74:64//HOMO SAPIENS (HUMAN).// P39192
  - F-PLACE2000061

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- F-PLACE2000062//GLUCOSE STARVATION-INDUCIBLE PROTEIN B (GENERAL STRESS PROTEIN B).//1.9e-06:108:37//BACILLUS SUBTILIS.//P26907
- 99 F-PLACE200072I/ZINC FINGER PROTEIN 165.//3.5e-34-175-49I/HOMO SAPIENS (HUMAN),/P49910 F-PLACE200097I/RIBONUCLEASE PANCREATIC (EC 3.1.27.5) (RNASE 1) (RNASE A).//0.36:39:38I/ONDAT-RA ZIBETHICUS (MUSKRAT),/P00081
  - F-PLACE2000100
  - F-PLACE2000103//TUBULIN ALPHA-4 CHAIN (FRAGMENTS).//0.18:32:37//ZEA MAYS (MAIZE).//P33626 F-PLACE2000111//CMRF35 ANTIGEN PRECURSOR.//0.056:107:27//HOMO SAPIENS (HUMAN).//Q08708
- F-PLACE200011/I/CMRF35 ANTIGEN PRECURSOR./I0.056·107:27/I/HOMO SAPIENS (HUMAN).//008708 F-PLACE2000115//DIAMINOPIMELATE EPIMERASE (EC 5.1.1.7) (DAP EPIMERASE) (FRAGMENT).//1.0:21: 52/CLOSTRIDIUM PERFRINGENS./I/Q46185
  - F-PLACE2000124//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.4e-37:108:68//HOMO SAPIENS (HU-MAN).//P39194
- 40 F-PLACE2000132//PROBABLE MEMBRANE ANTIGEN GP85.//0.99:133:29//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03224
  - F-PLACE2000136/NASOACTIVE INTESTINAL POLYPEPTIDE RECEPTOR 2 PRECURSOR (VIP-R-2) (PITUI-TAY ADENYLATE CYCLASE ACTIVATING POLYPEPTIDE TYPE III RECEPTOR) (PACAP TYPE III RECEP-TOR) (PACAP-R-3)//0.83:65:32/MUS MUSCULUS (MOUSE)://P41588
- 45 F-PLACE2000140

- F-PLACE2000164I/TIPD PROTEIN.//5.7e-12:190:28I/DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//
  - F-PLACE2000170//BACTERIOCIN CARNOBACTERIOCIN BM1 PRECURSOR (CARNOBACTERIOCIN B1).//
    1.0:30:26//CARNOBACTERIUM PISCICOLA //P38579
- F-PLACE2000172
  - F-PLACE2000176/IHYPOTHETICAL PROTEIN AF0526/I0.7644.43/IARCHAEOGLOBUS FULGIDUS //029724 F-PLACE2000187//EM-LIKE PROTEIN GEA6.//0.84.42:35//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS).// Q02973
  - E-DI ACE2000216
- F-PLACE2000223//NEUROTOXIN III (LQQ III),//0.99:38:34//LEIURUS QUINQUESTRIATUS QUINQUESTRIA-TUS (EGYPTIAN SCORPION).//P01487
  - F-PLACE2000235
    - F-PLACE2000246//RING CANAL PROTEIN (KELCH PROTEIN).//5.1e-37:121:42//DROSOPHILA MELA-

### NOGASTER (FRUIT FLY) //Q04652

- F-PLACE2000264//!!!! ALU SUBFAMILY SB2 WARNING ENTRY !!!!/2.4e-05:77:42//HOMO SAPIENS (HU-MAN).//P39191
- F-PLACE2000274//DYNEIN BETA CHAIN, CILIARY.//5.3e-46:232:45//TRIPNEUSTES GRATILLA (HAWAIAN SEA URCHIN) //P23098
  - F-PLACE2000302//TRICHOHYALIN.//1.5e-06:215:29//ORYCTOLAGUS CUNICULUS (RABBIT).//P37709
  - F-PLACE2000305//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.3e-06:33:66//HOMO SAPIENS (HUMAN).//
- F-PLACE2000317//TOXIN C13S1C1 PRECURSOR.//0.44:45:33//DENDROASPIS ANGUSTICEPS (EASTERN GREEN MAMBA).//P18329
  - F-PLACE2000335//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//7.9e-08:35:71//HOMO SAPIENS (HUMAN).//
  - F-PLACE2000341//SODIUM/GLUCOSE COTRANSPORTER 1 (NA(+)/GLUCOSE COTRANSPORTER 1) (HIGH AFFINITY SODIUM-GLUCOSE COTRANSPORTER).//0.014:141:24//ORYCTOLAGUS CUNICULUS (RABBIT).//
- 15 P11170 F-PLACE2000342/HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION./15.7e-09:96:38// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMIPV)./IP41479
  - F-PLACE2000347//ZINC FINGER PROTEIN 177.//5.9e-05:49:53//HOMO SAPIENS (HUMAN).//Q13360
  - F-PLACE2000359//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.5e-10:69:52//HOMO SAPIENS (HUMAN).// P39194
  - F-PLACE2000366

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- F-PLACE2000371//ATROPHIN-1 (DENTATORUBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN).//1.5e-05:216: 29//HOMO SAPIENS (HUMAN).//P54259
- F-PLACE2000373//MAX BINDING PROTEIN MNT (ROX PROTEIN) (MYC ANTAGONIST MNT).//0.27:63:33//
- 25 HOMO SAPIENS (HUMAN).//Q99583 F-PLACE2000379//HYPOTHETICAL GENE 1 PROTEIN.//0.72:120:31//EQUINE HERPESVIRUS TYPE 1
  - (STRAIN AB4P) (EHV-1).//P28978 F-PLACE2000394//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.95:40:42//HOMO SAPIENS (HUMAN).//
- 30 F-PLACE2000398//RIBONUCLEASE PRECURSOR (EC 3.1.27.-)//0.88:88:31//AEROMONAS HYDROPHILA.// Q07465
  - F-PLACE2000399/T-CELL SURFACE GLYCOPROTEIN E2 PRECURSOR (E2 ANTIGEN) (CD99) (MIC2 PRO-TEIN) (12E7).//7.6e-16:180:39/HOMO SAPIENS (HUMAN).//P14209
  - F-PLACE2000404//PROBABLE LEUCYL-TRNA SYNTHETASE (EC 6.1.1.4) (LEUCINE-TRNA LIGASE) (LEURS).//1.7e-94:243:64//CAENORHABDITIS ELEGANS.//Q09996
  - F-PLACE2000411//SERINE/THREONINE PROTEIN PHOSPHATASE 5 (EC 3.1.3.16) (PP5) (PROTEIN PHOS-PHATASE T) (PPT) (FRAGMENT) //1.2e-09:78:39//MUS MUSCULUS (MOUSE) //060676
  - F-PLACE2000419//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.6e-20:61:62//HOMO SAPIENS (HUMAN).// P39188
- 40 F-PLACE2000425//HYPOTHETICAL 11.9 KD PROTEIN IN MSB2-UGA1 INTERGENIC REGION.//0.98:75:32// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53211
  - F-PLACE2000427//INSULIN PRECURSOR.//0.98:55:34//CERCOPITHECUS AETHIOPS (GREEN MONKEY) (GRIVET).//P30407
- F-PLACE2000433//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//7.5e-07:65:50//HOMO SAPIENS (HUMAN).//
  45 P39188
  - F-PLACE2000435
    - F-PLACE2000438//HYPOTHETICAL 67.9 KD PROTEIN ZK688.8 IN CHROMOSOME III.//4.7e-66:178:47// CAENORHABDITIS ELEGANS.//P34678
  - F-PLACE2000450//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//2.1e-23:88:62//HOMO SAPIENS (HUMAN).// P39195
  - F-PLACE2000455/TOXIN II (TOXIN II.10.9.2) (FRAGMENT).//0.093:18:44//CENTRUROIDES LIMPIDUS LIMPIDUS (MEXICAN SCORPION).//P45630
  - F-PLACE2000458//CADHERIN-RELATED TUMOR SUPPRESSOR PRECURSOR (FAT PROTEIN).//3.1e-23: 165:40//DROSOPHILA MELANOGASTER (FRUIT FLY).//P33450
- 55 F-PLACE2000465//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.6e-23:73:63//HOMO SAPIENS (HUMAN).//
  - F-PLACE2000477///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/4.4e-37:90:78//HOMO SAPIENS (HUMAN).// P39194

- F-PLACE3000004//EYES ABSENT HOMOLOG 3.//1.1e-09:27:100//MUS MUSCULUS (MOUSE),//P97480
  F-PLACE3000009//PUTATIVE CUTICLE COLLAGEN C09G5.6.//0.0061:148:34//CAENORHABDITIS ELECANS.//OP0457
- F-PLACE3000020//ADENYLATE CYCLASE, OLFACTIVE TYPE (EC 4.6.1.1) (TYPE III) (ATP PYROPHOS-PHATE-LYASE) (ADENYLYL CYCLASE).//8.8e-93:193:92//RATTUS NORVEGICUS (RAT).//P21932
- F-PLACE3000229/f305 RIBOSOMAL PROTEIN L31E, I/O 15:50:38/METHANOCOCCUS JANNASCHII./IP54009 F-PLACE3000059/TCP1-CHAPERONIN COFACTOR A./IO 96:50:34/IBOS TAURUS (BOVINE)./IP48427 F-PLACE3000070/HYPOTHETICAL 17.1 KD PROTEIN IN PURS 3'REGION./IO 29:22:59/SACCHAROMYCES
  - F-PLACE3000070/HYPOTHETICAL 17.1 KD PROTEIN IN PUR5 3'REGION.//0.29:22:59//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38898
- F-PLACE3000103/LYSIS PROTEIN (E PROTEIN) (GPE).//0.99:53:32//BACTERIOPHAGE ALPHA-3.//P31280 F-PLACE3000119//!!! ALU SUBFAMILY SB WARNING ENTRY !!!!/5.4e-41:87:78//HOMO SAPIENS (HUMAN).// P39189
  - F-PLACE3000121//VESICULAR TRAFFIC CONTROL PROTEIN SEC151/1.0e-07:269:22//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P22224
- 15 F-PLACE3000124//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.2e-29:97:73//HOMO SAPIENS (HUMAN).// P39188
  - F-PLACE3000136//PARS INTERCEREBRALIS MAJOR PEPTIDE D1 (PMP-D1).//0.77:26:42//LOCUSTA MIGRA-TORIA (MIGRATORY LOCUST).//P80059
  - F-PLACE3000142//HYPOTHETICAL 7.1 KD PROTEIN IN NAD2 3'REGION (ORF 63).//0.82:34:41//MARCHAN-TIA POLYMORPHA (LIVERWORT).//P38468
  - F-PLACE3000145//TENSIN.//3.5e-91:238:74//GALLUS GALLUS (CHICKEN).//Q04205
    - F-PLACE3000147//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//4.4e-30:61:65//HOMO SAPIENS (HUMAN).//
- F-PLACE3000148//POL POLYPROTEIN [CONTAINS: PROTEASE (EC 3.4.23.-); REVERSE TRANSCRIPTASE

  (EC 2.7.7.49); ENDONUCLEASEL//1.4e-18:226:34//GIBBON APE LEUKEMIA VIRUS.//P21414
- F-PLACE3000155//EXTENSIN PRECURSOR (PROLINE-RICH GLYCOPROTEIN).//0.00014:107:33//ZEA MAYS (MAIZE).//P14918
- F-PLACE3000156//POL POLYPROTEIN [CONTAINS: PROTEASE (EC 3.4.23.-); REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE].//2.7e-19:169:30//BABOON ENDOGENOUS VIRUS (STRAIN M7).//P10272
- F-PLACE3000157//PROBABLE SERINE/THREONINE-PROTEIN KINASE CY50.16 (EC 2.7.1.-).//0.0061:92:30// MYCOBACTERIUM TUBERCULOSIS.//Q11053
  - F-PLACE3000158//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//5.7e-49:56:80//HOMO SAPIENS (HUMAN).// P39189
    - F-PLACE3000160//DNA TRANSFORMATION PROTEIN TFOX (COMPETENCE ACTIVATOR) (PROTEIN SXY).// 0.39:94:34//HAEMOPHILUS INFLUENZAE.//P43779
      - F-PLACE3000169//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/5.6e-28:99:59//HOMO SAPIENS (HUMAN).// P39193
    - F-PLACE3000194//PROLINE-RICH PROTEIN LAS17.//0.91:80:36//SACCHAROMYCES CEREVISIAE (BAK-ER'S YEAST).//Q12446
- 40 F-PLACE3000197//NEUROFILAMENT TRIPLET M PROTEIN (160 KD NEUROFILAMENT PROTEIN) (NF-M).// 0.24:119:32//GALLUS GALLUS (CHICKEN).//P16053
  - F-PLACE3000199//EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN).// 0.76:87:37//NICOTIANA TABACUM (COMMON TOBACCO).//P13983
- F-PLACE3000207//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.5e-09:32:78//HOMO SAPIENS (HUMAN).//
  45 P39188
- F-PLACE3000208

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- F-PLACE3000218//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.2e-34:96:70//HOMO SAPIENS (HUMAN).//
- F-PLACE3000220//OSTEOCALCIN (GAMMA-CARBOXYGLUTAMIC ACID-CONTAINING PROTEIN) (BONE GLA- PROTEIN) (BGP).//0.46:13:53//CANIS FAMILIARIS (DOG).//P81455
- F-PLACE3000221//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.8e-24:178:45//HOMO SAPIENS (HUMAN).//
  - F-PLACE3000226//30S RIBOSOMAL PROTEIN S18.//0.98:38:34//NEISSERIA GONORRHOEAE.//O07815
- F-PLACE3000230//METALLOTHIONEIN (MT).//0.97:25:48//OREOCHROMIS MOSSAMBICUS (MOZAMBIQUE TILAPIA) (TILAPIA MOSSAMBICA).//P52726
  - F-PLACE3000242//MELANOMA-ASSOCIATED ANTIGEN 8 (MAGE-8 ANTIGEN).//8.0e-21:121:39//HOMO SA-PIENS (HUMAN).//P43361
    - F-PLACE3000244//PROTEIN TSG24 (MEIOTIC CHECK POINT REGULATOR).//2.3e-125:264:87//MUS MUS-

- CULUS (MOUSE) //P53995
- F-PLACE3000254//RTOA PROTEIN (RATIO-A).//0.99:142:23//DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).//P54681
- F-PLACE3000271//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.2e-12:63:53//HOMO SAPIENS (HUMAN).//
  - F-PLACE3000276//COLLAGEN ALPHA 1(VIII) CHAIN PRECURSOR (ENDOTHELIAL COLLAGEN).//1.0:55:38//
    HOMO SAPIENS (HUMAN).//P27658
  - F-PLACE3000304//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.0028:31:54//HO-MO SAPIENS (HUMAN).//P30808
- F-PLACE3000310//ATROPHIN-1 (DENTATORUBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN).//0.98:82:34// RATTUS NORVEGICUS (RAT).//P54258
  - F-PLACE3000320
  - F-PLACE3000322//GLYCINE-RICH CELL WALL STRUCTURAL PROTEIN 1 PRECURSOR.//2.2e-22:61:52//
    ORYZA SATIVA (RICE).//P25074
- 15 F-PLACE3000331//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//0.32:15:53//HOMO SAPIENS (HUMAN).//P22532
  - F-PLACE3000339//CHORION PROTEIN S19.//0.34:89:37//DROSOPHILA VIRILIS (FRUIT FLY).//P24516 F-PLACE3000341//NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 1 (EC 1.6.5.3) (FRAGMENT).//1.0:47:38//
- COTURNIX COTURNIX APONICA (JAPANESE QUAIL).//P24968

  F-PLACE3000550/SERINETH/REONINE-PROTEIN KINASE SULU (EC 2.7.1.-).//3.9e-50:168:60//
  CAENORHABDITIS ELEGANS //P46549
  - F-PLACE3000352//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.8e-29:76:71//HOMO SAPIENS (HUMAN).//
- F-PLACE3000353//POLYPEPTTOE N-ACETYLGALACTOSAMINYLTRANSFERASE (EC 2.4.1.41) (PROTEIN-UDP ACETYLGALACTOSAMINYLTRANSFERASE) (UDP-GALNAC:POLYPEPTIDE, N- ACETYLGALACTOS-
- AMINYLTRANSFERASE) (GALNAC-T1)//3.0e-09:100:41/IH/DMO SAPIENS (HUMAN)//010472
  F-PLACE3000362/IHYPOTHETICAL PROTEIN TP0064/I1.0:75:26/I/TREPONEMA PALLIDUM//083103
  F-PLACE3000363/METALLOTHIONEIN (MT)//0.067:42:33//ASTACUS FLUVIATILIS (BROAD-FINGERED
- CRAYFISH) (ASTACUS ASTACUS //P55951

  F-PLACE3000365//LYSIS PROTEIN (E PROTEIN) (GPE)//1.0:65-27//BACTERIOPHAGE PH-K.//Q38040
  F-PLACE3000373//RETROVIRUS-RELATED ENV POLYPROTEIN//1.5e-18:90:47//HOMO SAPIENS (HU-
  - MAN).//P10267 F\_PLACE3000388
    - F-PLACE3000399//!!!ALU SUBFAMILY SP WARNING ENTRY !!!!//6.3e-45:60:75//HOMO SAPIENS (HUMAN).//
  - F-PLACE3000400

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- F-PLACE3000401//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//3.6e-09:46:73//HOMO SAPIENS (HUMAN).// P39188
- F-PLACE3000402//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.036:43:44//HOMO SAPIENS (HUMAN).//
  - F-PLACE3000405//POSTERIOR PITUITARY PEPTIDE.//0.70:25:40//BOS TAURUS (BOVINE),//P01154
    F-PLACE3000406//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//4.36-09:49:67//HOMO SAPIENS (HUMAN),//
- F-PLACE3000413//MALE SPECIFIC SPERM PROTEIN MST87F.//0.12:42:40//DROSOPHILA MELANOGASTER

  (FRUIT FLY).//P08175
  - F-PLACE3000416//CYLICIN I (MULTIPLE-BAND POLYPEPTIDE I).//0.67:236:21//BOS TAURUS (BOVINE).// P35662
    - F-PLACE3000425//PROLINE-RICH PEPTIDE P-B.//0.45:19:42//HOMO SAPIENS (HUMAN).//P02814
  - F-PLACE3000455//AMELOGENIN, CLASS I PRECURSOR./I0.0073:81:43//BOS TAURUS (BOVINE).//P02817 F-PLACE3000475//8.6 KD TRANSGLUTAMINASE SUBSTRATE.//10:53:32//TACHYPLEUS TRIDENTATUS (JAPANESE HORSESHOE GRABI.//P81281
    - F-PLACE3000477//MUSCARINIC TOXIN 7 (MT-7).//0.13:55:32//DENDROASPIS ANGUSTICEPS (EASTERN GREEN MAMBA) //PR0970
- F-PLACE400009//MYOSIN HEAVY CHAIN, SMOOTH MUSCLE ISOFORM (SMMHC) (FRAGMENT).//7.0e-19: 180:27//HOMO SAPIENS (HUMAN).//P35749
  - F-PLACE4000014//X-LINKED HELICASE II (X-LINKED NUCLEAR PROTEIN) (XNP).//3.2e-15:193:30//HOMO SAPIENS (HUMAN).//P46100
    - F-PLACE4000034//BRIDE OF SEVENLESS PROTEIN PRECURSOR.//0.0024:97:29//DROSOPHILA MELA-

NOGASTER (FRUIT FLY).//P22815

- F-PLACE4000049//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.8e-32:79:75//HOMO SAPIENS (HUMAN).// P39194
- F-PLACE4000052//ATP-BINDING CASSETTE TRANSPORTER 1.//2.2e-99:178:97//MUS MUSCULUS (MOUSE).//P41233
  - F-PLACE4000063//IMMEDIATE-EARLY PROTEIN.//0.0017:159:25//HERPESVIRUS SAIMIRI (STRAIN 11).// Q01042
  - F-PLACE4000089
- F-PLACE4000093
- 10 F-PLACE4000100/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.5e-14:68:60//HOMO SAPIENS (HUMAN).//
  - F-PLACE4000106//1A PROTEIN ICONTAINS: HELICASE: METHYLTRANSFERASEI.//1.0:46:41//BROAD BEAN MOTTLE VIRUS //Q00020
- F-PLACE4000128//HYPOTHETICAL PROTEIN E-115.//0.00020:101:30//HUMAN ADENOVIRUS TYPE 2.// 15 P03290
  - F-PLACE4000129//CORNIFIN B (SMALL PROLINE-RICH PROTEIN IB) (SPR-IB) (14.9 KD PANCORNULIN).// 0.15:57:31//HOMO SAPIENS (HUMAN).//P22528
    - F-PLACE4000131
    - F-PLACE4000147//COMPETENCE PHEROMONE PRECURSOR.//1.0:45:24/BACILLUS SUBTILIS.//P45453 F-PLACE4000156//ZINC FINGER PROTEIN 136.//2.1e-88:194:59//HOMO SAPIENS (HUMAN).//P52737
- 20 F-PLACE4000192//ZINC FINGER PROTEIN 142 (KIAA0236) (HA4654).//0.083:148:26//HOMO SAPIENS (HU-MAN).//P52746
  - F-PLACE4000211//CALPHOTIN.//0.20:43:39//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q02910
- F-PLACE4000222/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.1e-05:20:85//HOMO SAPIENS (HUMAN).// 25
  - F-PLACE4000230//DIHYDROFOLATE REDUCTASE (EC 1.5.1.3) / THYMIDYLATE SYNTHASE (EC 2.1.1.45) (DHFR-TS), I/1.0:96:28//TRYPANOSOMA BRUCEI BRUCEI, I/Q27783
  - F-PLACE4000233 F-PLACE4000247//METALLOTHIONEIN (MT).//1.0e-05:34:41//PLEURONECTES PLATESSA (PLAICE).//
- 30 F-PLACE4000250//VPU PROTEIN (ORF-X PROTEIN) (UPX PROTEIN).//0.99:33:42//CAPRINE ARTHRITIS EN-
  - CEPHALITIS VIRUS (CAEV).//P31834 F-PLACE4000252//MALE SPECIFIC SPERM PROTEIN MST84DB.//0.42:24:45//DROSOPHILA MELA-
  - NOGASTER (FRUIT FLY).//Q01643
- 35 F-PLACE4000259//PRE-MRNA SPLICING HELICASE BRR2 (EC 3.6.1.-),//3.5e-09:189:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32639
  - F-PLACE4000261//PEREGRIN (BR140 PROTEIN)://5.0e-11:103:37//HOMO SAPIENS (HUMAN)://P55201 F-PLACE4000269//INTRACELLULAR PROTEIN TRANSPORT PROTEIN USO1.//0.037:181:25//SACCHARO-
- MYCES CEREVISIAE (BAKER'S YEAST),//P25386 40 F-PLACE4000270//COAGULATION FACTOR VII PRECURSOR (EC 3.4.21.21).//1.0:46:39//MUS MUSCULUS
- (MOUSE).//P70375 F-PLACE4000300//50S RIBOSOMAL PROTEIN L32.//0.81:28:46//THERMUS AQUATICUS (SUBSP. THER-
  - MOPHILUS).//P80339 F-PLACE4000320//FKBP-RAPAMYCIN ASSOCIATED PROTEIN (FRAP) (RAPAMYCIN TARGET PROTEIN).//
- 45 1.6e-29:44:93//HOMO SAPIENS (HUMAN).//P42345 F-PLACE4000323
  - F-PLACE4000326//PARATHYMOSIN.//0.0018:54:48//HOMO SAPIENS (HUMAN).//P20962
  - F-PLACE4000344//EPIDERMAL GROWTH FACTOR (EGF) (FRAGMENT), I/O.97:28:42//SUS SCROFA (PIG), I/ Q00968
- F-PLACE4000367//NEUROTOXIN 1 (TOXIN SHP-I) (SHNA) (NEUROTOXIN SHI).//1.0:33:36//STOICHACTIS 50 HELIANTHUS (CARRIBEAN SEA ANEMONE) (STICHODACTYLA HELIANTHUS).//P19651
  - F-PLACE4000369//EXTENSIN\_PRECURSOR\_(PROLINE-RICH\_GLYCOPROTEIN).//0.071:42:42//SORGHUM VULGARE (SORGHUM).//P24152
- F-PLACE4000379//!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/1.4e-16:54:77//HOMO SAPIENS (HUMAN).//
  - F-PLACE4000387//PHOTOSYSTEM II 4 KD REACTION CENTRE PROTEIN PRECURSOR.//0.25:21:52//HOR-DEUM VULGARE (BARLEY), AND SECALE CEREALE (RYE), //P25877
    - F-PLACE4000392//FERROCHELATASE (EC 4.99.1.1) (PROTOHEME FERRO-LYASE) (HEME SYNTHETASE)

- (FRAGMENT) //0.91:36:50//YERSINIA PSEUDOTUBERCULOSIS //Q05338
- F-PLACE4000401//!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//4.4e-29:96:67//HOMO SAPIENS (HUMAN).//
- F-PLACE4000411///!!! ALU SUBFAMILY J WARNING ENTRY !!!!//2.3e-18:41:73//HOMO SAPIENS (HUMAN).//
  - F-PLACE4000431//PRE-MRNA SPLICING HELICASE BRR2 (EC 3.6.1.-).//5.4e-21:237:33//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32639
  - F-PLACE4000445//HYPOTHETICAL 99.7 KD PROTEIN IN SDL1 5'REGION PRECURSOR.//0.00081:210:26// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40442
- 10 F-PLACE4000450//TRANSCRIPTION FACTOR HBP-1A (HISTONE-SPECIFIC TRANSCRIPTION FACTOR HBP1).//0.020:87;33//TRITICUM AESTIVUM (WHEAT).//P23922
  - F-PLACE4000465//METALLOTHIONEIN-IL (MT-1L) (MT1X).//0.20:18:38//HOMO SAPIENS (HUMAN).//P80297 F-PLACE4000487//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.5e-19:73:52//HOMO SAPIENS (HUMAN).//
- 15 F-PLACE4000489
  - F-PLACE4000494//NPC DERIVED PROLINE RICH PROTEIN 1 (NDPP-1).//0.17:130:30//MUS MUSCULUS (MOUSE).//Q03173
    - F-PLACE4000521//RETROVIRUS-RELATED POL POLYPROTEIN [CONTAINS REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASE] (FRAGMENT).//3.0e-05:50:36//MUS MUSCULUS (MOUSE).//P10400
- F-PLACE4000522//NEUROGENIC LOCUS NOTCH HOMOLOG PROTEIN 1 PRECURSOR.//1.8e-45:231:47// RATTUS NORVEGICUS (RAT).//Q07008
  - F-PLACE4000548//CYTOCHROME C-551 (C551).//0.96:50:34//ECTOTHIORHODOSPIRA HALOPHILA.// P00122
- F-PLACE4000588//PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE FAF (EC 3.12.15) (UBIQUITIN TINIOLESTERASE FAF) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE FAF) (DEUBIQUITINATING ENZYME FAF) (FAT FACETS PROTEIN) //1 6e-28:223.36//DROSOPHILA MELANOGASTER (FRUIT FLY).//P55824 F-PLACE000581//P-SELECTIN PRECURSOR (GRANULE MEMBRANE PROTEIN 140) (GMP-140) (PADGEM) (CD62P) (LEUKOCYTE-ENDOTHELIAL CELL ADHESION MOLECULE 3) (LECAM3).//9.7e-11:166.281/HOMO SAPIENS (HUMAN)./P6109
- 30 F-PLACE4000590//POL POLYPROTEIN [CONTAINS: PROTEASE (EC 3.4.23.-); REVERSE TRANSCRIPTASE (EC 2.7.7.49); ENDONUCLEASEL//1.6e-17:134:35//GIBBON APE LEUKEMIA VIRUS.//P21414
  - F-PLACE4000593//GONADOTROPIN-RELEASING HORMONE RECEPTOR (GNRH-R).//1.0:54:29//RATTUS NORVEGICUS (RAT).//P30969
- F-PLACE4000612//GAG POLYPROTEIN [CONTAINS: CORE PROTEIN P15; INNER COAT PROTEIN P12; CORE SHELL PROTEIN P30].//2.6e-14:221:32//MOLONEY MURINE SARCOMA VIRUS (STRAIN TS110).// P272604
  - F-PLACE4000638//HYPOTHETICAL 9.3 KD PROTEIN IN NRDB-INAA INTERGENIC REGION.//0.65:37:40//ES-CHERICHIA COLI //P37910
- F-PLACE4000650//ZINC FINGER PROTEIN 16 (ZINC FINGER PROTEIN KOX9) (FRAGMENT).//1.0:33:33//HO-40 MO SAPIENS (HUMAN).//P17020
  - F-PLACE4000654

- F-PLACE4000670//HYPOTHETICAL 44.1 KD PROTEIN IN RPB5-CDC28 INTERGENIC REGION.//1.6e-07:161: 25//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P33313
- F-SKNMC1000011//PUTATIVE IMPORTIN BETA-4 SUBUNIT (KARYOPHERIN BETA-4 SUBUNIT).//7.4e-15:223: 45 31//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//060100
- F-SKNMC1000013//TRANSCRIPTION FACTOR BF-2 (BRAIN FACTOR 2) (BF2) (CBF-2) (T-14-6).//0.0013:128: 35//GALLUS GALLUS (CHICKEN).//Q98937
  - F-SKNMC1000046//CUTICLE COLLAGEN 1.//0.0010:154:33//CAENORHABDITIS ELEGANS.//P08124
  - F-SKNMC1000050//CALPAIN 2, LARGE [CATALYTIC] SUBUNIT (EC 3.4.22.17) (CALCIUM-ACTIVATED NEUTRAL PROTEINASE) (CANP) (M-TYPE).//3.2e-41:87:98//HOMO SAPIENS (HUMAN).//P17655
  - F-SKNMC1000091/NTAK PROTEIN (NEURAL-AND THYMUS-DERIVED ACTIVATOR FOR ERBB KINASES).// 0.0032:154:35//HOMO SAPIENS (HUMAN).//O14511
  - F-THYRO1000017//PUTATIVE PYRIDOXAMINE 5'-PHOSPHATE OXIDASE (EC 1.4.3.5) (PNP/PMP OXI-DASE).//1.6e-23:124:37//CAENORHABDITIS ELEGANS.//Q20939
- 55 F-THYRO1000026//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//3.0e-13:54:66//HOMO SAPIENS (HUMAN).//
  - F-THYRO1000034//HYPOTHETICAL 10.4 KD PROTEIN.//0.16:44:34//HEPATITIS B VIRUS (SUBTYPE AYW).// P03163

- F-THYRO1000035/CAMPATH-1 ANTIGEN PRECURSOR (CD52 ANTIGEN) (CDW52) (CAMBRIDGE PATHOL-OGY 1 ANTIGEN) // 0.83:59:37//MACACA FASCICULARIS (CRAB EATING MACAQUE) (CYNOMOLGUS MON-KEY) // P32763
- F-THYRO1000040//60S RIBOSOMAL PROTEIN L37 (FRAGMENT).//0.25:23:39//BOS TAURUS (BOVINE).// P79244
  - F-THYRO1000070//HYPOTHETICAL 29.3 KD PROTEIN (ORF92).//2.3e-11:133:36//ORGYIA PSEUDOTSUGA-TA MULTICAPSID POLYHEDROSIS VIRUS (OPMNPV).//O10341
  - F-THYRO1000072//C-PROTEIN, SKELETAL MUSCLE SLOW-ISOFORM.//1.5e-14:205:29//HOMO SAPIENS (HUMAN).//Q00872
- F-THYRO1000085

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- F-THYRO1000092//SPERM MITOCHONDRIAL CAPSULE SELENOPROTEIN (MCS).//0.063:59:33//HOMO SA-PIENS (HUMAN).//P49901
- F-THYRO1000107
- F-THYRO1000111//LINE-1 REVERSE TRANSCRIPTASE HOMOLOG://5.0e-58:110:67//NYCTICEBUS COU-CANG (SLOW LORIS)://P08548
  - F-THYRO1000121//SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SF3A66).//2.6e-06:134:35//MUS MUSCULUS (MOUSE).//Q62203
    - F-THYRO1000124//TÉNECIN 3 PRECURSOR.//0.047:76:35//TENEBRIO MOLITOR (YELLOW MEALWORM).//
- 20 F-THYRO1000129//FBROSIN (FRAGMENT).//0.35:43:34/MUS MUSCULUS (MOUSE).//Q60791
  - F-THYRO1000132/!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//8.7e-14:104:42//HOMO SAPIENS (HUMAN).// P39188
    - F-THYRO1000156
- F-THYRO1000163//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//3.7e-20:71:71//HOMO SAPIENS (HUMAN).//
- 30 P39192 F-THYRO1000187
  - F-THYRO1000190//PROTEIN TRANSPORT PROTEIN SEC61 BETA 2 SUBUNIT.//0.060:50:42//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P52871
  - F-THYRO1000197
- 35 F-THYRO1000199//HYPOTHETICAL 49.8 KD PROTEIN D2007.5 IN CHROMOSOME III.//2.0e-06:88:35// CAENORHABDITIS ELEGANS.//34379
  - F-THYRO1000206
  - F-THYRO1000221
  - F-THYRO1000241//HYPOTHETICAL 11.8 KD PROTEIN IN HE65-PK2 INTERGENIC REGION.//1.0:51:35// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV).//P41661
- F-THYRO1000242//ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2),//7.4e-37:137:36//HOMO SA-PIENS (HUMAN),//P51523
  - F-THYRO1000253//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.11:21:52//HO-MO SAPIENS (HUMAN) //P30808
- 45 F-THYRO1000270/WDNM1 PROTEIN PRECURSOR.//0.40:52:32//MUS MUSCULUS (MOUSE).//Q62477 F-THYRO1000279//BETA CRYSTALLIN A4.//0.97:64:26//BOS TAURUS (BOVINE).//P11842
  - F-THYRO1000288//POTENTIAL CAAX PRENYL PROTEASE 1 (EC 3.4.24.-) (PRENYL PROTEIN- SPECIFIC ENDOPROTEASE 1) (PPSEP 1).//3.4e-48:142:42//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//
- Q10071
  F-THYRO1000320//ZINC FINGER PROTEIN 14 (ZFP-14) (KROX-9 PROTEIN) (FRAGMENT).//0.87:35:45//MUS
  MUSCULUS (MOUSE)./P10755
  - F-THYRO1000327//HYPOTHETICAL 64.7 KD PROTEIN F26E4.11 IN CHROMOSOME I.//0.00010:75:26// CAENORHARDITIS FLEGANS //P90859
- F-THYRO1000343//CHROMOGRANIN A PRECURSOR (CGA) [CONTAINS: PANCREASTATIN; BETA-GRANIN; WE-141.//0.88:107:26//MUS MUSCULUS (MOUSE).//P26339
  - F-THYRO1000358//SELENIUM-BINDING LIVER PROTEIN.//4.6e-25:49:81//MUS MUSCULUS (MOUSE).// P17563
    - F-THYRO1000368//LOCOMOTION-RELATED PROTEIN HIKARU GENKI PRECURSOR.//1.0:136:26//DRO-

SOPHILA MELANOGASTER (FRUIT FLY).//Q09101

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- F-THYRO1000381//GAG POLYPROTEIN [CONTAINS: CORE PROTEIN P15; INNER COAT PROTEIN P12; CORE SHELL PROTEIN P30; NUCLEOPROTEIN P101.//0.032;99;35//SIMIAN SARCOMA VIRUS.//P03330
- GRAY SEAL.//P38592
- F-THYRO1000394//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.00019:48:37//HOMO SAPI-ENS (HUMAN).//P22531
- F-THYRO1000395//RING CANAL PROTEIN (KELCH PROTEIN).//1.2e-33:186:38//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- 10 F-THYRO1000401//50S RIBOSOMAL PROTEIN L7/L12 (FRAGMENT).//0.57:67:31//STAPHYLOCOCCUS AU-REUS.//P48860
  - F-THYRO1000438//ATP SYNTHASE PROTEIN 8 (EC 3.6.1.34) (A6L).//1.0:42:38//STRONGYLOCENTROTUS PURPURATUS (PURPLE SEA URCHIN).//P15997
  - F-THYRO1000452//BACTERIOCIN CARNOBACTERIOCIN A PRECURSOR (PISCICOLIN 61).//0.31:34:44// CARNOBACTERIUM PISCICOLA.//P38578
  - F-THYRO1000471//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2,1e-31:94:72//HOMO SAPIENS (HUMAN).// P39194
    - F-THYRO1000484//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//5.9e-08:30:86//HOMO SAPIENS (HUMAN).// P39195
- 20 F-THYRO1000488//EARLY NODULIN 55-2 PRECURSOR (N-55-2) (NODULIN-315).//0.93:98:27//GLYCINE MAX (SOYBEAN).//Q02917
  - F-THYRO1000501//DOWN REGULATORY PROTEIN OF INTERLEUKIN 2 RECEPTOR.//2.4e-51:198:50//MUS MUSCULUS (MOUSE).//P15533
- F-THYRO1000502//HUNCHBACK PROTEIN (FRAGMENT).//0.84:41:43//APIS MELLIFERA (HONEYBEE).//
  P31504
  - F-THYRO1000505//HYPOTHETICAL BHLF1 PROTEIN.//0.99:231:33//EPSTEIN-BARR VIRUS (STRAIN B95-8)
    (HUMAN HERPESVIRUS 4).//P03181
  - F-THYRO1000558//ANTITHROMBIN-III PRECURSOR (ATIII) (FRAGMENT).//0.47:58:37//GALLUS GALLUS (CHICKEN).//Q03352
- 30 F-THYRO1000569//COLLAGEN ALPHA 1(I) CHAIN (FRAGMENTS).//0.00048:64:42//RATTUS NORVEGICUS (RAT).//P02454
  - F-THYRO1000570//HYPOTHETICAL 11.6 KD PROTEIN IN ACS1-GCV3 INTERGENIC REGION.//0.94:61:32// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P39725
- F-THYRO1000585//SPLICING FACTOR, ARGININE/SERINE-RICH 6 (PRE-MRNA SPLICING FACTOR SRP55).//0.050:104:36//HOMO SAPIENS (HUMAN).//Q13247
  - F-THYRO1000596/INFECTED CELL PROTEIN ICP34.5 (NEUROVIRULENCE FACTOR ICP34.5),//0.99:37:40// HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN MGH-10),//P37319 F-THYRO1000602/IFAMZP30-47 PROTEIN (FRAGMENT),//0.88:61:34//EIMERIA ACERVULINA.//P21959
- F-THYRO1006052/EAR31 / F32583

  F-THYRO100605//SUPPRESSOR PROTEIN SRP40.//0.0016:116:26//SACCHAROMYCES CEREVISIAE

  40 (BAKER'S YEAST) / F32583
  - F-THYRO1000625//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//3.4e-33:88:78//HOMO SAPIENS (HUMAN).//
- F-THYRO1000637//METALLOTHIONEIN A (MT A).//1.0:23:43//SPARUS AURATA (GILTHEAD SEA BREAM).//
- 45 F-THYRO1000641//PHOTOSYSTEM II 10 KD PHOSPHOPROTEIN.//0.99:26:46//CYANIDIUM CALDARIUM (GALDIERIA SULPHURARIA) //0.19925
  - F-THYRO1000658/IIII ALU SUBFAMILY SB WARNING ENTRY IIII/I1.5e-49:116:69/HOMO SAPIENS (HU-MAN).//P39189
  - F-THYRO1000662//DNA-DAMAGE-INDUCIBLE PROTEIN P.//3.7e-15:119.43//ESCHERICHIA COLI.//Q47155 F-THYRO1000666//KINESIN-LIKE PROTEIN KLP1.//1.0e-44:232:41//CHLAMYDOMONAS REINHARDTII.// PAR870
    - F-THYRO1000676//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//2.1e-15:144:39//HOMO SAPIENS (HU-MAN) //P39193
- F-THYRO1000684//HYPOTHETICAL 73.5 KD PROTEIN IN SCS3-RPS2 INTERGENIC REGION.//0.00033:84:

  35 30//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53129
- F-THYRO1000699/!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//0.97:20:85//HOMO SAPIENS (HUMAN).//
  - F-THYRO1000712//!!! ALU SUBFAMILY J WARNING ENTRY !!!!//4.2e-10:69:59//HOMO SAPIENS (HUMAN).//

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- F-THYRO1000715//SALIVARY PROLINE-RICH PROTEIN PRECURSOR (CLONES CP3, CP4 AND CP5) [CON-TAINS: BASIC PEPTIDE IB-6; PEPTIDE P-H].//4.6e-10:204:32//HOMO SAPIENS (HUMAN).//P04280 E-THYPO1000734
- F-THYRO1000748//HYPOTHETICAL PROTEIN KIAA0411 (FRAGMENT).//1.8e-46:130:70//HOMO SAPIENS (HUMAN).//043295
  - F-THYRO1000756//ALPHA-N-ACETYLGALACTOSAMINIDE ALPHA-2,6-SIALYLTRANSFERASE (EC 2.4.99.-) (ST6GALNACIII) (STY).//1.1e-06:95:31//RATTUS NORVEGICUS (RAT).//Q64686
  - F-THYRO1000777//CUTICLE COLLAGEN 2C (FRAGMENT).//0.0031:119:34//HAEMONCHUS CONTORTUS.//
  - F-THYRO1000783//MYOSIN IC HEAVY CHAIN.//0.0014:121:37//ACANTHAMOEBA CASTELLANII (AMOEBA).//
  - F-THYRO1000787//HUNCHBACK PROTEIN (FRAGMENT),//0.54:25:52//PHOLCUS PHALANGIOIDES.//
- F-THYRO1000793//PRE-MRNA SPLICING FACTOR PRP9.//0.91:3 0:36//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P19736
- F-THYRO1000796

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- F-THYRO1000805//HYPOTHETICAL 7.3 KD PROTEIN IN 100 KD PROTEIN REGION.//0.081:31:38//HUMAN ADENOVIRUS TYPE 41.//P23691
- 20 F-THYRO1000815//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//6.0e-30:81:70//HOMO SAPIENS (HUMAN).// P39195
  - F-THYRO1000829//NEUROTOXIN III (BOM III).//0.022:32:34//BUTHUS OCCITANUS MARDOCHEI (MOROC-CAN SCORPION) //P13488
- F-THYRO 1000843//HYPOTHETICAL 7.7 KD PROTEIN IN GENES 5-4 INTERGENIC REGION (ORF 109).//0.98: 25:44//BACTERIOPHAGE P22.//P26750
  - F-THYRO1000852//SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185),//7.3e-09:83:42//VOLVOX CART-FRI //P21997
  - F-THYRO1000855//ANTIFREEZE PEPTIDE 4 PRECURSOR.//1.0:54:35//PSEUDOPLEURONECTA AMERICA-NUS (WINTER FLOUNDER) //P02734
- 30 F-THYRO1000865/!!!! ALU SUBFAMILY J WARNING ENTRY!!!!/5.2e-17:66:57//HOMO SAPIENS (HUMAN).// P30188
  - F-THYRO1000895//!!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//1.0e-12:58:62//HOMO SAPIENS (HUMAN).// P39189
    - F-THYRO1000916///!!! ALU SUBFAMILY SB WARNING ENTRY !!!!//2.0e-32:101:69//HOMO SAPIENS (HU-MAN).//P39189
    - F-THYRO1000926//NITROGEN FIXATION REGULATORY PROTEIN.//5.5e-05:108:27//KLEBSIELLA OXYTO-CA.//P56267
    - F-THYRO1000934//PYRROLINE-5-CARBOXYLATE REDUCTASE (EC 1.5.1.2) (P5CR) (P5C REDUCTASE).// 3.9e-50:147:40//HOMO SAPIENS (HUMAN).//P32322
- 40 F-THYRO1000951//DIHYDROXYACETONE KINASE (EC 2.7.1.29) (GLYCERONE KINASE).//1.8e-31:136:56// CITROBACTER FREUNDII.//P45510
  - F-THYRO1000952//HYPOTHETICAL 182.0 KD PROTEIN IN NMD5-HOM6 INTERGENIC REGION.//2.4e-05:91: 34//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P47170
- F-THYRO1000974//MITOCHONDRIAL ATP-DEPENDENT RNA HELICASE SUV3 PRECURSOR.//1.0:35:40//
  SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32580
- F-THYRO1000975
  - F-THYRO1000983//HYPOTHETICAL 48.1 KD PROTEIN B0403.2 IN CHROMOSOME X.//1.3e-20:96:51//
    CAENORHABDITIS ELEGANS.//Q11076
  - F-THYRO1000984//GTP-BINDING ADP-RIBOSYLATION FACTOR HOMOLOG 1 PROTEIN.//0.011:76:34//DRO-SOPHILA MELANOGASTER (FRUIT FLY).//P25160
- F-THYRO1000988
  - F-THYRO1001003//HYPOTHETICAL 8.1 KD PROTEIN IN MSCL-RPLQ INTERGENIC REGION.//0.97:60:31// ESCHERICHIA COLI.//P36675
- F-THYRO1001031//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!//9.5e-18:56:66//HOMO SAPIENS (HUMAN).//
  55 P39195
  - F-THYRO1001033//TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521.//5.0e-13:126:35//HOMO SAPI-ENS (HUMAN).//P31948
    - F-THYRO1001062//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.1e-35:97:79//HOMO SAPIENS (HUMAN).//

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- F-THYRO1001093//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//6.4e-13:70:57//HOMO SAPIENS (HUMAN).//
- F-THYRO1001100//ZINC FINGER X-LINKED PROTEIN ZXDA (FRAGMENT).//4.2e-63:219:63//HOMO SAPIENS (HUMAN).//P98168
  - F-THYRO1001120//SPLICEOSOME ASSOCIATED PROTEIN 49 (SAP 49) (SF3B53).//0.00068:160:31//HOMO SAPIENS (HUMAN) //015427
- F-THYRO1001121/VERY HYPOTHETICAL 20.6 KD PROTEIN C56F8.15 IN CHROMOSOME I.//0.37:158:28// SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10263
- 10 F-THYRO1001133//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!/7.3e-15:59:66/HOMO SAPIENS (HUMAN).// P39188 F-THYRO1001134//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE M) [CONTAINS: PEPTIDE P-D] (FRAG-
  - MENT).//0.00088:159:29//HOMO SAPIENS (HUMAN).//P10161 F-THYRO1001142////III ALU SUBFAMILY SQ WARNING ENTRY I/!!//3.0e-29:81:71//HOMO SAPIENS (HUMAN).//
- F-THYRO1001173//CYTOCHROME C OXIDASE POLYPEPTIDE VIIS (EC 1.9.3.1).//0.88:51:35//DICTYOSTEL-IUM DISCOIDEUM (SLIME MOLD).//P20610
  - F-THYRO1001177//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//3.0e-24:91:68//HOMO SAPIENS (HUMAN).//
- 20 F-THYRO1001189//MKR2 PROTEIN (ZINC FINGER PROTEIN 2).//7.3e-27:165:39//MUS MUSCULUS (MOUSE).//P08043
  - F-THYRÓ1001204//BASIC PROLINE-RICH PEPTIDE P-E (IB-9).//0.67:42:42//HOMO SAPIENS (HUMAN).//
- F-THYRO1001213///!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//2.9e-16:61:68//HOMO SAPIENS (HUMM).// P39194
- F-THYRO1001262//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.4e-36:50:84//HOMO SAPIENS (HUMAN).//
- F-THYRO1001271//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.62: 126:30//STREPTO-MYCES FRADIAE.//P20186
- 30 F-THYRO1001287//HYPOTHETICAL 91.2 KD PROTEIN IN RPS4B-SCH9 INTERGENIC REGION.//1.9e-26:208: 37//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38888
  - F-THYRO1001290//GIANT HEMOGLÓBIN AIV CHAIŃ (FRAGMENT).//1.0:31:38//LAMELLIBRACHIA SP. (DEEP-SEA GIANT TUBE WORM).//P20413
    - F-THYRO1001313//VACUOLAR PROTEIN SORTING-ASSOCIATED PROTEIN VPS5.//0.00042:105:31//SAC-CHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q92331
  - F-THYRO1001320//COLLAGEN ALPHA 1(III) CHAIN.//0.27:57:38//BOS TAURUS (BOVINE).//P04258
    - F-THYRO1001321//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//5.5e-20:74:64//HOMO SAPIENS (HUMAN).// P39188
- F-THYRO1001322//HYPOTHETICAL 7.2 KD PROTEIN.//0.66:49:30//VACCINIA VIRUS (STRAIN COPENHA-49 GEN).//P21123
  - F-THYRO1001347//TOXIN F-VIII PRECURSOR (TOXIN TA2) (TOXIN DAF8).//0.94:61:36//DENDROASPIS AN-GUSTICEPS (EASTERN GREEN MAMBA).//P01404
  - F-THYRO1001363//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//0.0025:23:73//HOMO SAPIENS (HUMAN).// P39188
- 45 F-THYRO1001365/MERSACIDIN PRECURSOR.//0.35:38:42//BACILLUS SP. (STRAIN HIL-Y85/54728).//
  - F-THYRO1001374//PROTEIN VDLD.//1.6e- 3:140:31//HELICOBACTER PYLORI (CAMPYLOBACTER PYLORI).//005729
  - F-THYRO1001401///!!! ALU SUBFAMILY SC WARNING ENTRY !!!!//0.047:43:48//HOMO SAPIENS (HUMAN).// P39192
- F-THYRO1001403

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- F-THYRO1001405//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.0068:26:42//HOMO SAPIENS (HUMAN) //P22531
- F-THYRO1001406//PUTATIVE STEROID DEHYDROGENASE KIK-I (EC 1.1.1.-).//3.1e-81:97:83//MUS MUSCU-LUS (MOUSE).//O70503
  - F-THYRO1001411//!!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.9e-26:89:74//HOMO SAPIENS (HUMAN).//
    - F-THYRO1001426//!!! ALU SUBFAMILY SP WARNING ENTRY !!!!//1.4e-09:55:61//HOMO SAPIENS (HUMAN).//

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- F-THYRO1001434//BETA-DEFENSIN 4 PRECURSOR (BNDB-4).//0.68:44:34//BOS TAURUS (BOVINE).//
- F-THYRO1001458//MYOSIN HEAVY CHAIN, NONMUSCLE TYPE B (CELLULAR MYOSIN HEAVY CHAIN, TYPE B) (NMMHC-B),//3.8e-64:216:62//HOMO SAPIENS (HUMAN),//P35580
  - F-THYRO1001480//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!/9.3e-29:88:75//HOMO SAPIENS (HUMAN).// P39194
    - F-THYRO1001487//HOMEOBOX PROTEIN HOX-B4 (HOX-2.6).//0.99:59:37//MUS MUSCULUS (MOUSE).// P10284
- 10 F-THYRO1001534//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//1.4e-14:40:82//HOMO SAPIENS (HUMAN).// P39194
  - F-THYRO1001537//HYPOTHETICAL 33.8 KD PROTEIN IN TWT1-FLO5 INTERGENIC REGION.//2.4e-07:142: 32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38892
- F-THYRO1001541///!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.98:26:61//HOMO SAPIENS (HUMAN).//
  P39195
- F-THYRO1001559//PROTEIN Q300 //2.6e-05:20:75//MUS MUSCULUS (MQUSE).//Q02722
  - F-THYRO1001570

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- F-THYRO1001573//SPERM MITOCHONDRIAL CAPSULE SELENOPROTEIN (MCS).//0.033:71:36//MUS MUS-CULUS (MOUSE).//P15265
- F-THYRO1001584//SUPPRESSOR PROTEIN SRP40.//2.1e-05:188:27//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P32583
  - F-THYRO1001595//RAS SUPPRESSOR PROTEIN 1 (RSU-1) (RSP-1 PROTEIN) (RSP-1).//6.1e-21:35:91//HO-MO SAPIENS (HUMAN).//Q15404
- F-THYRO1001602//TRK SYSTEM POTASSIUM UPTAKE PROTEIN TRKH.//1.0:57:42//HAEMOPHILUS INFLU-ENZAE.//P44843
- F-THYRO1001605//VENOM BASIC PROTEASE INHIBITORS IX AND VIIIB.//1.0:34:38//BUNGARUS FASCIA-TUS (BANDED KRAIT).//P25660
  - F-THYRO1001617//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.0e-18:55:81//HOMO SAPIENS (HUMAN).// P39194
- 30 F-THYRO1001637//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.00020:25:80//HOMO SAPIENS (HU-MAN).//P39195
  - F-THYRO1001656//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//0.0091:54:42//MUS MUSCULUS (MOUSE).//P05142
  - F-THYRO1001661//HYPOTHETICAL 21.1 KD PROTEIN IN SSR-SERA INTERGENIC REGION (O182).//0.033: 77:35//ESCHERICHIA COLI.//P09160
  - F-THYRO1001671//(2'-5')OLIGOADENYLATE SYNTHETASE 1 (EC 2.7.7.-) ((2-5')OLIGO(A) SYNTHETASE 1) (2-5A SYNTHETASE 1) (P46/P41) (E18/E16).//4.3e-34:207:34//HOMO SAPIENS (HUMAN).//P00973
  - F-THYRO1001673//!!!!ÄLU SUBFAMILY SQ WARNING ENTRY !!!!//3.9e-08:49:65//HOMO SAPIENS (HUMAN).// P39194
- 40 F-THYRO1001703//HYPOTHETICAL 69.8 KD PROTEIN IN BDF1-SFP1 INTERGENIC REGION.//6.4e-16:134: 35//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q06053 F-THYRO1001706
  - F-THYRO1001721//RING CANAL PROTEIN (KELCH PROTEIN).//2.7e-27:191:36//DROSOPHILA MELA-NOGASTER (FRUIT FLY).//Q04652
- 45 F-THYRO1001738/MATING PROCESS PROTEIN MID2 (SERINE-RICH PROTEIN SMS1) (PROTEIN KINASE A INTERFERENCE PROTEIN)./I0.0032:105:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P36027 F-THYRO1001745
  - F-THYRO1001746//GENE 10 PROTEIN.//1.0:55:30//SPIROPLASMA VIRUS SPV1-R8A2 B.//P15901
  - F-THYRO1001772//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//1.2e-05:41:63//HOMO SAPIENS (HUMAN).// P39188
  - F-THYRO1001793//HYPOTHETICAL 21.6 KD PROTEIN F37A4.2 IN CHROMOSOME III.//1.5e-26:161:42//CAENORHABDITIS ELEGANS.//P41880
    - F-THYRO1001809//LATENCY-RELATED PROTEIN 2.//0.49:74:27//HERPES SIMPLEX VIRUS (TYPE 1 / STRAIN F).//P17589
- F-THYRO1001828//PROTEINASE INHIBITOR.//0.11:34:50//SOLANUM MELONGENA (EGGPLANT) (AUBER-GINE).//P01078
  - F-THYRO1001854//ACYL-COA-BINDING PROTEIN HOMOLOG (ACBP) (DIAZEPAM BINDING INHIBITOR HO-MOLOG) (DBI).//0.63:50:38//RANA RIDIBUNDA (LAUGHING FROG) (MARSH FROG).//P45883

- F-THYRO1001895//!!!! ALU SUBFAMILY J WARNING ENTRY !!!!//6.1e-09:72:47//HOMO SAPIENS (HUMAN).// P39188
- F-THYRO1001907//TRYPOMASTIGOTE DECAY-ACCELERATING FACTOR (T-DAF) (FRAGMENT).//0.79:36: 44//TRYPANOSOMA CRUZL//026327
- F-VESEN1001122/IHOMEOBOX PROTEIN HB9.//0.57:84:32/IHOMO SAPIENS (HUMAN) //P50219
   F-Y79A41000013//METALLOTHIONEIN B (MT-B).//0.034:35:48//SALMO SALAR (ATLANTIC SALMON).// P57720
  - F-Y79AA1000033//CHOLECYSTOKININ.//0.97:49:30//PSEUDEMYS SCRIPTA (SLIDER TURTLE).//P80345
- F-Y79AA1000037//DNA-BINDING PROTEIN BMI-1.//1.4e-23-80:60//HOMO SAPIENS (HUMAN).//P35226
  F-Y79AA1000059//HYPOTHETICAL 35.5 KD PROTEIN IN TRANSPOSON TN4556.//0.0075:127:36//STREPTOMYCFS FRADIAF (197016)
  - F-Y79AA1000065//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.022:135:29//HOMO SAPIENS (HUMAN).//P10162
- F-Y79AA1000131//REGULATORY PROTEIN E2://1.1e-05:175:26//HUMAN PAPILLOMAVIRUS TYPE 24://
  15 P50770
- F-Y79AA1000181//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//1.4e-06:187:29//MUS MUSCULUS (MOUSE) //P05143
  - F-Y79AA1000202//HYPOTHETICAL PROLINE-RICH PROTEIN (FRAGMENT).//6.2e-09:47:53//OWENIA FUSI-FORMIS //P21260
- F-Y79AA1000214//HISTONE H2A VARIANT.//1.7e-50-107:100I/GALLUS GALLUS (CHICKEN).//P02272 F-Y79AA1000230I/GONADOLIBERIN I PRECURSOR (LHRH I) (LUTEINIZING HORMONE RELEASING HORMONE I) (GONADOTROPIN RELEASING HORMONE I) (GNRH I) (LULIBERIN I).//0.27:64:34//HOMO SAPIENS (HUMAN).//P01148
- F-Y79AA1000231//HYPOTHETICAL 47.9 KD PROTEIN M021B04.12.//2.5e-72:277:53//ARABIDOPSIS THAL-IANA (MOUSE-EAR CRESS).//004658
  - F-Y79AA1000258//PROLINE-RICH PROTEIN MP-2 PRECURSOR.//2.8e-08:174:35//MUS MUSCULUS
  - F-Y79AA1000268//COLLAGEN ALPHA 1(III) CHAIN (FRAGMENT).//0.00020:176:33//RATTUS NORVEGICUS (RAT).//P13941
- 30 F-Y79AA1000313//HYPOTHETICAL 54.0 KD PROTEIN C32A3.1 IN CHROMOSOME III.//0.092:127:21// CAENORHABDITIS ELEGANS.//Q09260
  - F-Y79AA1000328//SEL-10 PROTEIN.//5.3e-05:129-28//CAENORHABDITIS ELEGANS.//Q93794
    F-Y79AA100034/IKERATIN, ULTRA HIGH-SULFUR MATRIX PROTEIN (UHS KERATIN).//1.0:73:30//OVIS AR-IES (SHEEP).//P26372
- 35 F-Y79AA1000346//COATOMER GAMMA SUBUNIT (GAMMA-COAT PROTEIN) (GAMMA-COP).//1.8e-95:205: 83/BOS TAURUS (BOVINE).//P53620
  - F-Y79AA1000349/IANTIFREEZE PEPTIDE 4 PRECURSOR.//0.036:37:54//PSEUDOPLEURONECTA AMERI-CANUS (WINTER FLOUNDER).//P02734
- F-Y79AA1000355//HYPOTHETICAL 18.2 KD PROTEIN ZK632.13 IN CHROMOSOME III.//0.0031:106:28//
  49 CAENORHABDITIS ELEGANS.//O10120
- F-Y79AA1000368//REDUCED VIABILITY UPON STARVATION PROTEIN 161.//1.4e-16:208:28//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P25343
- F-Y79AA1000405//LIGHT-HARVESTING PROTEIN B-800-850, ALPHA CHAIN C (ANTENNA PIGMENT PRO-TEIN, ALPHA CHAIN C) (LH II-C ALPHA).//0.98:50:30//RHODOPSEUDOMONAS PALUSTRIS.//P35103
- 45 F-Y79AA1000410/!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!//7.9e-20:62:79//HOMO SAPIENS (HUMAN).// P39 194 F-Y79AA1000420//HYPOTHETICAL 27.7 KD PROTEIN IN UME3-HDA1 INTERGENIC REGION.//1.4e-06:86:38//
  - SACCHAROMYCES CEREVISIAE (BAKER'S YEAST), I/P53970
    F-Y79AA1000469//HYPOTHETICAL 48.4 KD PROTEIN F44B9.5 IN CHROMOSOME III.//2.8e-34:211:40//
- 50 CAENORHABDITIS ELESANS.//P34426 F-Y79A41000480//HYPOTHETICAL 63.2 KD PROTEIN C1F3.09 IN CHROMOSOME 1.//3.9e-15:90:32//
  - SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10414 F-779AA1000538//!!!! ALU SUBFAMILY SX WARNING ENTRY!!!!//0.37:41:48//HOMO SAPIENS (HUMAN).//
- F-Y79AA1000539//SPLICING FACTOR, ARGININE/SERINE-RICH 4 (PRE-MRNA SPLICING FACTOR SRP75).// 1.8e-21:190:37//HOMO SAPIENS (HUMAN).//Q08170
  - F-Y79AA1000540//SPERM PROTAMINE P1.//0.00045:66:45//DASYURUS VIVERRINUS (SOUTHEASTERN QUOLL), AND DASYURUS HALLUCATUS.//P42135

- F-Y79A-41000560//ALPHA-ADAPTIN C (CLATHRIN ASSEMBLY PROTEIN COMPLEX 2 ALPHA-C LARGE CHAIN) (100 KD COATED VESICLE PROTEIN C) (PLASMA MEMBRANE ADAPTOR HA2/AP2 ADAPTIN ALPHA C SUBUNIT).//1.6e-79:186:87//MUS MUSCULUS (MOUSE)
- F-Y79AA1000574//AKLAVINONE C-11 HYDROXYLASE (EC 1.-.-.) (FRAGMENT).//0.010:35:60//STREPTOMY-CES PELICETIUS //P32009
- F-Y79AA1000589//32.3 KD PROTEIN IN CWP1-MBR1 INTERGENIC REGION.//4.5e-27:197:36//SACCHARO-MYCES CEREVISIAE (BAKER'S YEAST).//P28320
- F-Y79AA1000627//ZINC FINGER PROTEIN 134.//1.6e-34:191:35//HOMO SAPIENS (HUMAN).//P52741
- F-Y79AA1000705//HYPOTHETICAL 128.5 KD HELICASE IN ATS1-TPD3 INTERGENIC REGION.//8.7e-36:250: 40//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P31380
  - F-Y79AA1000734//PEROXISOMAL MEMBRANE PROTEIN PMP30A (PMP31) (PEROXIN-11A).//0.00037:108: 27//CANDIDA BOIDINII (YEAST).//Q00316
  - F-Y79AA1000748//HYPOTHETICAL 61.3 KD PROTEIN F25B5.5 IN CHROMOSOME III./1.0e-23:210:34// CAENORHABDITIS ELEGANS.//Q09316
- 15 F-Y79AA1000752//PUTATIVE HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN X (HNRNP X) (CBP).// 1.4e-53:156:68//MUS MUSCULUS (MOUSE).//Q61990
  - F-Y79AA1000774//HYPOTHETICAL 77.9 KD PROTEIN IN RRN10-MCM2 INTERGENIC REGION.//1.2e-11:231: 26//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38205
  - F-Y79AA1000782//CUTICLE COLLAGEN 2.//0.012:56:35//CAENORHABDITIS ELEGANS.//P17656
- 20 F-Y79AA1000784//HISTIDINE-RICH GLYCOPROTEIN PRECURSOR.//1.3e-08:82:39//PLASMODIUM LOPHU-RAE.//P04929
  - F-Y79AA1000794//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS).//0.043:13:53//HO-MO SAPIENS (HUMAN).//P30808
- F-Y79AA1000800//PRIA PROTEIN PRECURSOR.//0.031:94:34//LENTINULA EDODES (SHIITAKE MUSH-PROOM) (LENTINUS EDODES) //001200
  - F-Y79AA1000802//HYPOTHETICAL 67.4 KD PROTEIN IN RPS3-PSD1 INTERGENIC REGION.//0.26:186:23// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53882
  - F-Y79AA1000805//AMP DEAMINASE (EC 3.5.4.6) (MYOADENYLATE DEAMINASE).//0.99:78:35//SCHIZOSAC-CHAROMYCES POMBE (FISSION YEAST).//P50998
- 30 F-Y79AA1000824//HYPOTHETICAL 81.7 KD PROTEIN IN MOL1-NAT2 INTERGENIC REGION.//3.4e-44:111: 49//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P48234
- F-Y79AA1000827//HYPOTHETICAL BHLF1 PROTEIN.//0.0046:187:33//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4)./P03181
  - F-Y79AA1000833//TUBULIN ALPHA-1 CHAIN.//1.0e-75:239:66//CRICETULUS GRISEUS (CHINESE HAM-STER).//P05209

- F-Y79AA1000850//SMALL PROLINE-RICH PROTEIN II (SPR-II) (CLONE 174N).//0.0078:57:31//HOMO SAPI-ENS (HUMAN).//P22532
- F-Y79AA1000962//MYOSIN HEAVY CHAIN, GIZZARD SMOOTH MUSCLE.//8.5e-11:241:26//GALLUS GALLUS (CHICKEN).//P10587
- 40 F-Y79AA1000966//ATP SYNTHASE A CHAIN (EC 3.6.1.34) (PROTEIN 6) .//0.69:122:31//TRYPANOSOMA BRUCEI BRUCEI.//P24499
  - F-Y79AA1000968//TRANSLATION INITIATION FACTOR EIF-2B GAMMA SUBUNIT (EIF-2B GDP-GTP EX-CHANGE FACTOR).//3.3e-102:211:93//RATTUS NORVEGICUS (RAT).//P70541
- F-Y79AA1000969//PROCOLLAGEN ALPHA 1(I) CHAIN PRECURSOR.//1.0:67:38//GALLUS GALLUS (CHICK-45 EN)://P02457
  - F-Y79AA1000976//INVOLUCRIN.//0.99:66:31//CEBUS ALBIFRONS (WHITE-FRONTED CAPUCHIN).//P24709 F-Y79AA1000985//PERICENTRIN.//1.1e-24:116:59//MUS MUSCULUS (MOUSE).//P48725
    - F-Y79AA1001023//HYPOTHETICAL 105.9 KD PROTEIN IN AAC3-RFC5 INTERGENIC REGION.//0.37:79:27// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38250
- 59 F-Y79AA1001041//SPERMATID-SPECIFIC PROTEIN T1 (CONTAINS: SPERM PROTAMINE SP1),///0.93:43:39// SEPIA OFFICINALIS (COMMON CUTTLEFISH).///P80001 F-Y79AA1001048//ACYL-COA DEHYDROGENASE. VERY-LONG-CHAIN SPECIFIC PRECURSOR (EC
  - 1.3.99.-) (VLCAD).//1.5e-51:211:52//BOS TAURUS (BOVINE).//P48818
- F-Y79AA1001061//!!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!.//3.8e-25:85:69//HOMO SAPIENS (HUMAN).//
  55 P39194
  - F-Y79AA1001068//PROCOLLAGEN ALPHA 1(II) CHAIN PRECURSOR [CONTAINS: CHONDROCALCIN].// 0.0015/207:33//MUS MUSCULUS (MOUSE).//P28481
  - F-Y79AA1001077//ADULT-SPECIFIC RIGID CUTICULAR PROTEIN 11.9 (ACP 11.9).//0.99:36:41//ARANEUS DI-

ADEMATUS (SPIDER) //P80515

F-Y79AA1001078//HYPOTHETICAL 88.1 KD PROTEIN K02D10.1 IN CHROMOSOME III.//1.0e-06:197:23// CAENORHABDITIS ELEGANS //P34492

F-Y79AA1001105//HOMEOBOX PROTEIN OTX2.//2.9e-62:163:79//MUS MUSCULUS (MOUSE).//P80206

F-Y79AA1001145//!!!! ALU SUBFAMILY SX WARNING ENTRY !!!!//0.024:42:59//HOMO SAPIENS (HUMAN).// P39195

F-Y79AA1001167//HYPOTHETICAL 7.1 KD PROTEIN IN IAP2-VLF1 INTERGENIC REGION.//0.96:20:50// AUTOGRAPHA CALIFORNICA NUCLEAR POLYHEDROSIS VIRUS (ACMNPV),//P41471

F-Y79AA1001177/JHYPOTHETICAL BHLF1 PROTEIN.//3.9e-05:135:34//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181

F-Y79AA1001185//PUTATIVE CUTICLE COLLAGEN C09G5.5.//0.00017:93:38//CAENORHABDITIS ELE-GANS//009456

F-Y79AA1001211 F-Y79AA1001216//TENSIN://0.012:134:32//GALLUS GALLUS (CHICKEN).//Q04205

15 F-Y79AA1001228//MUCIN 2 PRECURSOR (INTESTINAL MUCIN 2).//0.088:75:34//HOMO SAPIENS (HUMAN).//

F-Y79AA1001233//ESTRADIOL 17 BETA-DEHYDROGENASE 1 (EC 1.1.1.62) (17-BETA-HSD 1) (17-BETA-HYDROXYSTEROID DEHYDROGENASE 1).//1.1e-40:139:51//RATTUS NORVEGICUS (RAT).//P51657

F-Y79AA1001236//HYPOTHETICAL 34.7 KD PROTEIN IN ORC2-TIP1 INTERGENIC REGION.//2.0e-22:108:53// SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P38238

F-Y79AA1001281

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F-Y79AA1001299//PROLINE-RICH PROTEIN MP-3 (FRAGMENT).//0.0022:49:44//MUS MUSCULUS (MOUSE).//P05143

F-Y79AA1001312//50S RIBOSOMAL PROTEIN L24, CHLOROPLAST PRECURSOR.//0.98:117:25//ARABIDOP-SIS THALIANA (MOUSE-EAR CRESS).//P92959

SIS I HALLIANA (MUUSE-EAR CRESS),//PS24999
F-Y79AA1001323//CORNIFIN (SMALL PROLINE-RICH PROTEIN I) (SPR-I) (SMALL PROLINE-RICH SQUA-MOUS CELL MARKER) (SPRP)///0.082-44-40//SUS SCROFA (PIG).//P35323

F-779A4101384/IAPOLIPOPROTEIN C-III PRECURSOR (APO-CIII)/I/0.99:47:40//MUS MUSCULUS (MOUSE).//P33622

30 F-Y79AA1001391//HOMEOBOX PROTEIN HOX-A13 (HOX-1J).//9.8e-58:157:62//HOMO SAPIENS (HUMAN).// P31271

F-Y79AA1001394//TRICHOHYALIN.//4.7e-08:121:36//HOMO SAPIENS (HUMAN).//Q07283

F-Y79AA1001402//ETS-DOMAIN TRANSCRIPTION FACTOR ERF.//0.0087:81:33//MUS MUSCULUS (MOUSE).//P70459

F-Y79AA1001493//HYPOTHETICAL 48.1 KD PROTEIN B0403.2 IN CHROMOSOME X.//4.5e-21:125:44//
CAENORHABDITIS ELEGANS.//Q11076

F-Y79AA1001511//HYPOTHETICAL 86.6 KD PROTEIN IN PFK1-TDS4 INTERGENIC REGION.//2.3e-17:249: 31//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P53313

F-YY9AA1001533//DNA-DIRECTED RNA POLYMERASE I49 KD POLYPEPTIDE (EC 2.7.7.6) (A49).//0.0089: 40 155:23//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//Q01080 F-YY9AA1001541

F-Y79AA1001548//!!!! ALU SUBFAMILY SC WARNING ENTRY !!!!!//1.1e-17:53:83//HOMO SAPIENS (HUMAN).// P39192

F-Y79AA1001555//MAJOR SURFACE ANTIGEN.//0.046:62:29//HEPATITIS B VIRUS.//P31873

45 F-Y79AA1001581//ACETYL-COENZYME A SYNTHETASE (EC 6.2.1.1) (ACETATE—COA LIGASE) (ACYL- AC-TIVATING ENZYME).//8.6e-11:144:31//ESCHERICHIA COLI.//P27550

F-Y79AA1001585//SPERM MITOCHONDRIAL CAPSULE SELENOPROTEIN (MCS).//0.012:64:40//MUS MUS-CULUS (MOUSE).//P15265

F-Y79AA1001594//CORNIFIN BETA.//0.61:88:31//MUS MUSCULUS (MOUSE).//O09116

50 F-Y79AA1001603//TRANSCRIPTION INITIATION FACTOR TFIID 135 KD SUBUNIT (TAFII-135) (TAFII-135) (TAFII-130), (TAFII-130

F-Y79AA1001613//ZINC FINGER PROTEIN 42 (MYELOID ZINC FINGER 1) (MZF-1).//4.5e-09:136:27//HOMO SAPIENS (HUMAN).//P28698

F-Y79AA1001647//HYPOTHETICAL 23.1 KD PROTEIN CY277.20C.//0.093:94:26//MYCOBACTERIUM TUBER-CULOSIS.//P71779

F-Y79AA1001665//HOMEOBOX PROTEIN DLX-2 (HOMEOBOX PROTEIN TES-1).//0.79:90:26//MUS MUSCU-LUS (MOUSE).//P40764

F-Y79AA1001679//LAMBDA-CRYSTALLIN.//1.6e-95:224:81//ORYCTOLAGUS CUNICULUS (RABBIT).//P14755

- F-Y79AA1001692//GERM\_CELL-LESS\_PROTEIN.//3.5e-08:78:38//DROSOPHILA\_MELANOGASTER\_(FRUIT FLY) //Q01820
- F-Y79AA1001696//INSULIN.//1.0:33:27//ANGUILLA ROSTRATA (AMERICAN EEL).//P42633
- F-Y79AA1001705//HYPOTHETICAL BHLF1 PROTEIN.//0.0013:192:33//EPSTEIN-BARR VIRUS (STRAIN B95-8) (HUMAN HERPESVIRUS 4).//P03181
- F-Y79AA1001711//PARATHYMOSIN (ZINC-BINDING 11.5 KD PROTEIN).//0.032;38;34//RATTUS NORVEGI-CUS (RAT) //P04550
  - F-Y79AA1001781

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- F-Y79AA1001805/VASODILATOR-STIMULATED PHOSPHOPROTEIN (VASP).//0.0063;128;30//HOMO SAPI-ENS (HUMAN).//P50552
  - F-Y79AA1001827//SPERM PROTAMINE P1.//0.015:45:40//DIDELPHIS MARSUPIALIS VIRGINIANA (NORTH AMERICAN OPOSSUM), AND MONODELPHIS DOMESTICA (SHORT-TAILED GREY OPOSSUM), //P35305 F-Y79AA1001846//!!! ALU SUBFAMILY J WARNING ENTRY!!!//2.4e-09:42:73//HOMO SAPIENS (HUMAN).// P39188
- 15 F-Y79AA1001848//KRUEPPEL PROTEIN (FRAGMENT), //1.8e-10:63:44//PSYCHODA CINEREA. //Q02035 F-Y79AA1001866//ZINC FINGER PROTEIN 90 (ZFP-90) (ZINC FINGER PROTEIN NK10)://0.00036:108:37// MUS MUSCULUS (MOUSE) //O61967
  - F-Y79AA1001874//OX40L RECEPTOR PRECURSOR (ACT35 ANTIGEN) (TAX-TRANSCRIPTIONALLY ACTI-VATED GLYCOPROTEIN-1 RECEPTOR) (CD134 ANTIGEN).//3.2e-07:100:35//HOMO SAPIENS (HUMAN).// P43489
  - F-Y79AA1001875//B-CELL GROWTH FACTOR PRECURSOR (BCGF-12 KD).//0.020:25:64//HOMO SAPIENS (HUMAN).//P20931
    - F-Y79AA1001923//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE K) [CONTAINS: PEPTIDE P-D] (FRAG-MENT),//0.016:83:36//HOMO SAPIENS (HUMAN),//P10162
- F-Y79AA1001963//PUTATIVE PRE-MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE SPAC10F6.02C.//8.1e-13:94:47//SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//O42643 F-Y79AA1002027//UBIQUITIN-CONJUGATING ENZYME E2-18 KD (EC 6.3.2.19) (UBIQUITIN- PROTEIN LIGASE) (UBIQUITIN CARRIER PROTEIN) (PM42).//9.8e-39:143:52//ARABIDOPSIS THALIANA (MOUSE-EAR CRESS) //P42743
- 30 F-Y79AA1002083//DNA-BINDING P52/P100 COMPLEX, 100 KD SUBUNIT (FRAGMENTS),//0.036:53:45//HO-MO SAPIENS (HUMAN).//P30808
  - F-Y79AA1002089//HYPOTHETICAL 49.1 KD PROTEIN F02A9.4 IN CHROMOSOME III.//0.12:171:22// CAENORHARDITIS ELEGANS //P34384
  - F-Y79AA1002093//MAX PROTEIN.//3.1e-07:111:29//BRACHYDANIO RERIO (ZEBRAFISH) (ZEBRA DANIO).// P52161
    - F-Y79AA1002103//SHORT NEUROTOXIN C.//0.040:21:47//AIPYSURUS LAEVIS (OLIVE SEA SNAKE).//
  - F-Y79AA1002115//HYPOTHETICAL PROTEIN MJ0827 //0 84:68:30//METHANOCOCCUS JANNASCHII // O58237
- 40 F-Y79AA1002125//HYPOTHETICAL 24.7 KD PROTEIN IN POM152-REC114 INTERGENIC REGION //3.4e-29: 197:39//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//P40206
  - F-Y79AA1002139//DNAJ PROTEIN HOMOLOG 1 (DROJ1).//1.9e-19:120:45//DROSOPHILA MELANOGASTER (FRUIT FLY).//Q24133
- F-Y79AA1002204//TBX6 PROTEIN (T-BOX PROTEIN 6),//0.0011:162:32//MUS MUSCULUS (MOUSE),//P70327 45 F-Y79AA1002208//ANKYRIN.//2.9e-08:231:29//MUS MUSCULUS (MOUSE).//Q02357
  - F-Y79AA1002209//TYROSYL-TRNA SYNTHETASE, MITOCHONDRIAL PRECURSOR (EC 6.1.1.1) (TYRO-SINE--TRNA LIGASE) (TYRRS).//3.7e-23:170:32//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST).//
- F-Y79AA1002210//CORNIFIN A (SMALL PROLINE-RICH PROTEIN IA) (SPR-IA) (SPRK) //0.0061:69:31//HOMO 50 SAPIENS (HUMAN).//P35321
  - F-Y79AA1002211//!!! ALU SUBFAMILY SP WARNING ENTRY !!!!/9.2e-10:43:62//HOMO SAPIENS (HUMAN).// P39193
  - F-Y79AA1002220
- F-Y79AA1002229/HYPOTHETICAL 60.7 KD PROTEIN C56F8.17C IN CHROMOSOME I.//1.9e-21:147:40// 55 SCHIZOSACCHAROMYCES POMBE (FISSION YEAST).//Q10264 F-Y79AA1002234
  - F-Y79AA1002246//MYOSIN IC HEAVY CHAIN.//0.00066:131:34//ACANTHAMOEBA CASTELLANII (AMOE-BA).//P10569

- F-Y79AA1002258//HYPOTHETICAL 103.9 KD PROTEIN ZK370.3 IN CHROMOSOME III.//4.3e-45:164:48// CAENORHABDITIS ELEGANS //002328
- F-Y79AA1002298//SALIVARY PROLINE-RICH PROTEIN PO (ALLELE M) [CONTAINS: PEPTIDE P-D] (FRAG-MENT).//0.0063:99:31//HOMO SAPIENS (HUMAN).//P10161
- F-Y79AA1002307

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- F-Y79AA1002311//HYPOTHETICAL 105.3 KD PROTEIN C01G6.5 IN CHROMOSOME III.//0.75:198:24// CAFNORHARDITIS FLEGANS.//P46012
- F-Y79AA1002351//CUTICLE COLLAGEN 34.//0.74:128:35//CAENORHABDITIS ELEGANS.//P34687 F-Y79AA1002361//GLC7-INTERACTING PROTEIN 2.//0.050:71:29//SACCHAROMYCES CEREVISIAE (BAK-
- ER'S YEAST).//P40036
  F-Y79AA1002399///NEUROMODULIN (AXONAL MEMBRANE PROTEIN GAP-43) (PP46) (B-50) (PROTEIN F1)
- (CALMODULIN-BINDING PROTEIN P-57)//1.088:30//CARASSIUS AURATUS (GOLDFISH),//P17691 F-Y79AA1002407//HYPOTHETICAL 31.5 KD PROTEIN IN YGP1-YCK2 INTERGENIC REGION //3.7e-16:232: 28//SACCHAROMYCES CEREVISIAE (BAKER'S YEAST),//P53899
- 15 F-Y79AA1002416//CTP SYNTHASE (EC 6.3.4.2) (UTP-AMMONIA LIGASE) (CTP SYNTHETASE).//6.7e-72: 162:84//HOMO SAPIENS (HUMAN).//P17812
  - F-Y79AA1002431//SMALL PROLINE RICH PROTEIN II (SPR-II) (CLONE 930).//0.81:34:41//HOMO SAPIENS (HUMAN).//P22531
  - F-Y79AA1002433//CELL DIVISION CONTROL PROTEIN 68.//0.00024:85:27//SACCHAROMYCES CEREVI-SIAE (BAKER'S YEAST).//P32558
    - F-Y79AA1002472//ZINC FINGER PROTEIN 35 (ZFP-35).//2.3e-60:217:44//MUS MUSCULUS (MOUSE).// P15620
- F-Y79A41002482//ZINC FINGER PROTEIN 141.//2.Oe-31:90:56/IHOMO SAPIENS (HUMAN).//Q15928
  F-Y79A41002487//HYPOTHETICAL 67.1 KD TRP-ASP REPEATS CONTAINING PROTEIN C57A10.05C IN
  CHROMOSOME I./10.18:41:38/ISCHIZOSACCHAROMYCES POMBE (FISSION YEASTI.//P87053

Homology Search Result Data 2.

[0300] The result of the homology search of the GenBank using the clone sequence of 5'-end except EST and STS.

[0301] Data include

the name of clone

definition of the top hit data.

the P-value: the length of the compared sequence: identity (%), and

the Accession No. of the top hit data, as in the order separated by II.

[0302] Data are not shown for the clones in which the P-value was higher than 1.

F-HEMBA1000005//Mouse tumor cell dnaJ-like protein 1 mRNA, complete cds.//3.4e-106:695:86//L16953

F-HEMBA1000012//Caenorhabditis-elegans cosmid C16C10, complete sequence.//1.5e-24:374:66//Z46787

- 40 F-HEMBA1000020//Homo sapiens beta 2 gene.//3.5e-112:529:90//X02344 F-HEMBA1000030//Rattus norvegicus G protein-coupled receptor kinase-associated ADP ribosylation factor GT-
  - Pase-activating protein (GIT1) mRNA, complete cds.//5.6e-124:743:88//AF085693
    F-HEMBA1000042//Human Chromosome 15q26.1 PAC clone pDJ460g16, WORKING DRAFT SEQUENCE, 3
- unordered pieces.//1.1e-25:529:65//AC004581

  F-HEMBA1000046//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 125/3, WORKING
  - DRAFT SEQUENCE I/3.2e-11:330.63/IAL033528
    F-HEMBA1000050//Homo sapiens DNA sequence from PAC 172K10 on chromosome 6q24. Contains STS, GSS and chromosome 6 fragment. complete sequence I/0.32:407:59/IAL022477
    - F-HEMBA1000076//Homo sapiens full-length insert cDNA clone ZB97G06.//6.2e-135:594:98//AF086182
    - F-HEMBA1000111//CIT-HSP-2291M18.TF CIT-HSP Homo sapiens genomic clone 2291M18 genomic survey sequence.//2.8e-16:132:79//AQ004134
      - F-HEMBA1000129//Homo sapiens chromosome 17, clone HCIT48C15, complete sequence.//8.6e-98:230:93// AC003104
    - F-HEMBA1000141//Homo sapiens mRNA for KIAA0797 protein, partial cds.//2.1e-167;791;98//AB018340
- F-HEMBA1000150//Homo sapiens mRNA for K/AA0788 protein, partial cds.//2.2e-44:242:96//AB018331 F-HEMBA1000156//Rattus norvegicus scaffold attachment factor B mRNA, complete cds.//1.1e-10.409:60// AFIRSR324
  - F-HEMBA1000158//Homo sapiens CAGH44 mRNA, partial cds.//1.6e-35:365:73//U80741

F-HEMBA1000168//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 321D2, WORKING DRAFT SEQUENCE //0.99:290:61//AL031033

F-HEMBA1000180//rat u2 small nuclear rna gene and flanks.//3.7e-18:112:98//K00034

F-HEMBA1000185

F-HEMBA1000193//Human FMR1 gene, 5' end.//0.0012:191:67//L19476

F-HEMBA1000201//Human Ini1 mRNA, complete cds.//2.0e-73:440:92//U04847

F-HEMBA1000213//Plasmodium falciparum MAL3P7, complete sequence.//0.90:332:59//AL034559

F-HEMBA1000216//Mus musculus hypoxia inducible factor three alpha mRNA, complete cds.//4.8e-117:585:83// AF060194

F-HEMBA1000227/H.sapiens CpG island DNA genomic Mse1 fragment, clone 179h6, reverse read cpg179h6.rt1a.//1,9e-14:95:98/Z64921

F-HEMBA1000231//H. sapiens CpG island DNA genomic Mse1 fragment, clone 90a5, reverse read cpg90a5.rt1a.// 5.1e.34:186:97//756144

F-HEMBA1000243//Human DNA sequence from PAC 440021 on chromosome X contains ESTs and STS.//4.1e-67:291:82//Z84481

F-HEMBA1000244//M.musculus Ank-1 mRNA for erythroid ankydn.//0.029:316:59//X69065

F-HEMBA1000251//Homo sapiens PAC clone DJ0988L12 from 7q11.23-q21.1, complete sequence.//0.35:467:60//AC004454

F-HEMBA1000264

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F-HEMBA1000280//Homo sapiens clone DJ0292L20, WORKING DRAFT SEQUENCE, 2 unordered pieces.//8.9e-20:218:78//AC004825

F-HEMBA1000282//Homo sapiens chromosome Y, clone 264,M,20, complete sequence.//4.2e-08:134:77//

F-HEMBA1000288//345L5.TPB CIT978SKA1 Homo sapiens genomic clone A-345L05, genomic survey sequence //1 16-06:152-73//817459

F-HEMBA1000290//Human ornithine decarboxylase gene, complete cds.//3.2e-11:507:62//M33764

F-HEMBA1000302//CIT-HSP-2169N13.TF CIT-HSP Homo sapiens genomic clone 2169N13, genomic survey sequence.//5.4e-06:86:88//B90730

F-HEMBA1000303/Mus musculus Plenty of SH3s (POSH) mRNA, complete cds. //f.9e-1117.701:88(/kF030131 F-HEMBA1000304//HS\_3006\_A1\_A09\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3006 Col=17 Row=A, genomic survey sequence //f.S.2e-40:24092//AQ118225

F-HEMBA1000307//Mus musculus mRNA for CDV-1R protein.//7.9e-127:815:84//Y10495

F-HEMBA1000327//HS\_3124\_B2\_H08\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3124 Col=16 Row=P, genomic survey sequence.//1.4e-11:87:96//AQ187492

35 F-HEMBA1000333

F-HEMBA 1000338//Homo sapiens chromosome X, PAC 671D9, complete sequence.//4.0e-66:271:84//AF031078 F-HEMBA 1000351//Homo sapiens PAC clone DJ0649P17 from 7q11.23-q21, complete sequence.//0.64:334:60// AC004848

F-HEMBA1000355//Pseudorabies virus serine/threonine kinase (ULPK) gene, partial cds and alkaline nuclease (AN) gene, complete cds://0.017:313:63//U25056

F-HEMBA1000356/i/Oryctolagus cuniculus troponin T cardiac isoform mRNA, 3" end of cds.//0.87:198:61/L40178 F-HEMBA1000357/i/HS\_3194\_A1\_D05\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3194 Cot=9 Row=6, genomic survey sequence //6.5e-90/436:98/i/AQ173748

F-HEMBA1000366//HS\_3027\_B2\_G06\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3027 Col=12 Row=N, genomic survey seguence.//0.0074:192:64//AQ128843

F-HEMBA 1000369//Human DNA sequence from clone 1039K5 on chromosome 22q12.3-13.2 Contains gene simliar to PICK1 perinuclear binding protein, gene similiar to monocarboxylate transporter (MCT3), ESTs, STS, GSS and a CpG island, complete sequence, VI 42-106:133.99//AL031587

F-HEMBA1000376//Homo sapiens chromosome 19, BAC CIT-B-393i15 (BC301323), complete sequence.//1.6e-22:659:63//AC006116

F-HEMBA1000387//Homo sapiens chromosome 12p13.3 clone RPCI11-264F23, WORKING DRAFT SE-OUENCE: 90 unordered pieces //3 2e-06:136:75//AC006122

F-HEMBA1000390//Homo sapiens BAC clone RG119C02 from 7p15, complete sequence.//3.5e-111:284:95// AC004520

55 F-HEMBA1000392//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 173D1, WORKING DRAFT SEQUENCE.//1.8e-39:332:80//AL031984

F-HEMBA1000396//Human Xq13 3' end of PAC 92E23 containing the X inactivation transcipt (XIST) gene, complete sequence.//9.5e-35:364:73//U80460

- F-HEMBA1000411//Human Xp22 contig of 3 PACS (R7-39D12, R7-134G1, R7-185L21) from the Roswell Park Cancer Institute, complete sequence.//8.1e-18:424:64//U96409
- F-HEMBA1000418//Drosophila melanogaster Oregon-R mitochondrial A+T region.//0.0026:564:59//U11584
- F-HEMBA1000422//Human DNA from chromosome 19 specific cosmid R30292, genomic sequence, complete sequence //9.2e-14:232:70//AC003112
- F-HEMBA1000428//Homo sapiens Xp22 BAC GSHB-590J6 (Genome Systems Human BAC library) complete sequence.//3.8e-37:408:69//AC004554
  - F-HEMBA1000434//Caenorhabditis elegans cosmid Y48E1B, complete sequence.//0.73:454:57//Z93393
- F-HEMBA1000442

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- F-HEMBA1000456//RPCI11-30J5.TV RPCI-11 Homo sapiens genomic clone RPCI-11-30J5, genomic survey sequence //6 3e-06/62/96//B85188
  - F-HEMBA1000459//Mus musculus hemin-sensitive initiation factor 2 alpha kinase mRNA, complete cds.//6.8e-70: 580:79//AF028808
- F-HEMBA1000460//Homo sapiens PAC clone DJ0593H12 from 7p31, complete sequence.//2.8e-154:746:98//
  AC004839
- F-HEMBA1000464//Homo saplens, clone hRPK.15\_A\_1, complete sequence.//4.8e-25:397:72//AC006213
  - F-HEMBA1000469//CIT-HSP-2167P21.TF CIT-HSP Homo sapiens genomic clone 2167P21, genomic survey sequence.//4.0e-83:406:99//B94160
  - F-HEMBA1000488//Homo sapiens Chromosome 22q11.2 PAC Clone p\_m11 in BCRL2-GGT Region, complete sequence.//4.2e-53:312:93//AC004033
  - F-HEMBA1000490//Campylobacter jejuni groES, groEL genes.//0.59:451:62//Y13334
    - F-HEMBA1000491/Murine sarcoma virus (Harvey-strain) H-ras transforming p21 gene.//8.6e-06:338:58//X00740 F-HEMBA1000501/Homo sapiens chromosome 17, clone hRPK.264\_B\_14, complete sequence.//9.4e-41:591:69//AC005884
- F-HEMBA1000504//Homo sapiens mRNA for osteoblast specific factor 2 (OSF-2os).//4.0e-07:57:100//D13666 F-HEMBA1000505
  - F-HEMBA1000508//\*\*\* SEQUENCING IN PROGRESS \*\*\* Homo sapiens chromosome 4, BAC clone C0135005; HTGS phase 1, WORKING DRAFT SEQUENCE, 23 unordered pieces.//0.035:329:61//AC004661
- F-HEMBA1000518//Caenorhabditis elegans cosmid C17H12.//0.96:425:58//AF045642

  F-HEMBA1000519//Homo sapiens Xp22 BAC GSHB-536K7 (Genome Systems Human BAC library) complete sequence //1.6e-53.300.89/(AC004616
- F-HEMBA1000520/Homo sapiens clone DJ0813F11, WORKING DRAFT SEQUENCE, 5 unordered pieces.//1.7e10:117:86/JAC006006
  - 10:117:86//AC006006 F-HEMBA1000523
- F-HEMBA1000531//Mus musculus Hsp70-related NST-1 (hsr.1) mRNA, complete cds.//3.9e-35:290:80//U08215 F-HEMBA1000534//Homo sapiens chromosome 17, clone hRPK.177\_H\_5, WORKING DRAFT SEQUENCE, 2 ordered pieces // 17-a5-36:28.71/JC.005973
  - F-HEMBA1000540//Arabidopsis thaliana DNA chromosome 4, BAC clone F7K2 (ESSAII project).//0.057:265:63//
- 40 F-HEMBA1000542/iRattus norvegicus mRNA for dipeptidyl peptidase III, complete cds.//1.2e-110:572:88/D89340 F-HEMBA1000545/Human DNA from cosmid L27h9, Huntington's Disease Region, chromosome 4p16.3 contains CGG island //7.5e-1378.08/9/2495.
  - F-HEMBA1000555//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 134 O19, WORKING DRAFT SEQUENCE.//3.2e-175:838:98//AL034555
- 45 F-HEMBA1000557//CIT-HSP-2369F15.TF CIT-HSP Homo sapiens genomic clone 2369F15, genomic survey sequence.//2.8e-32:315:78//AQ074611
  - F-HEMBA1000561//Rattus norvegicus Olf-1/EBF associated Zn finger protein Roaz mRNA, alternatively spliced form, complete cds.//3.4e-69:665:72//U92564
  - F-HEMBA1000563//Plasmodium falciparum 3D7 chromosome 12 PFYAC1383 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered pieces.//0.59:261:61//AC005504
- F-HEMBA1000568//HS\_3243\_B2\_A12\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3243 Col=24 Row=B, genomic survey sequence.//3.1e-54:323:91//AQ219628
  - F-HEMBA1000569//M.musculus mRNA for GPI-anchored protein.//1.4e-19:440:61//X89571
    F-HEMBA1000575//Plasmodium falciparum 3D7 chromosome 12 PFYAC357 genomic sequence, WORKING
- 55 DRAFT SEQUENCE, 7 unordered pieces,//0.0016;55f:57//AC005506
  F-HEMBA1000588//Mus musculus FLI-LRR associated protein-1 mRNA, complete cds://1.7e-11:132:79//
  - AF045573
  - F-HEMBA1000591//Homo sapiens mRNA for E1B-55kDa-associated protein.//7.3e-43:228:97//AJ007509

- F-HEMBA1000592//Mus musculus clone OST7314, genomic survey sequence.//7.3e-07:68:94//AF046733 F-HEMBA1000594//Human DNA sequence from PAC 306D1 on chromosome X contains ESTs.//8.7e-71:553.79// 783872
- F-HEMBA1000604//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 237J2, WORKING DRAFT SEQUENCE.//2.9e-21:158:75//AL021394
- F-HEMBA1000608//Homo sapiens mRNA for KIAA0456 protein, partial cds.//1.1e-118:561:99//AB007925
  - F-HEMBA1000622//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-152E5, complete sequence.//2.2e-28:428:70//AC004382
  - F-HEMBA1000636//Human CpG island sequence, clone Q28B8.//1.0e-15;274;68//D85773
- F-HEMBA1000637//Homo sapiens mRNA for KIAA0690 protein, partial cds.//6.7e-137:639:99//AB014590 F-HEMBA1000655//, complete sequence.//5.1e-83:685:80//AC005815
  - F-HEMBA1000657//Rattus norvegicus ADP-ribosylation factor-directed GTPase activating protein mRNA, complete cds //1.1e-91:597:84/IU35776
  - F-HEMBA1000662//Homo sapiens clone DJ0853H20, WORKING DRAFT SEQUENCE, 5 unordered pieces.// 0.019:695:57//AC004907
- 15 0.019:R95.57/IAC004907 F-HEMBA1000673/I/Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 229A8, WORKING DRAFT SEQUENCE //1.5e-48:325:85/IZ86090
  - F-HEMBA1000682//Homo sapiens (subclone 5 g5 from P1 H25) DNA sequence.//7.7e-61:615:74//L43411
  - F-HEMBA1000686
- 20 F-HEMBA1000702
  - F-HEMBA1000705//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-628 genomic sequence, WORKING DRAFT SEQUENCE, 9 unordered pleces.//0.0037;569:57//AC005507
  - F-HEMBA1000719//Streptomyces coelicolor cosmid 1C2://2.0e-09:483:62//AL031124
  - F-HEMBA1000722//Toxoplasma gondii chloroplast, complete genome //0.00058:762:57//U87145
- 25 F-HEMBA1000726//H.sapiens HLA-DRB1\*15 gene.//9.8e-49:189:89//X88791
  - F-HEMBA1000727//CIT-HSP-387P22.TRB CIT-HSP Homo sapiens genomic clone 387P22, genomic survey sequence.//0.0054;206;67//B60158
  - F-HEMBA1000747
- F-HEMBA1000749/Human DNA sequence from clone 522P13 on chromosome 6p21.31-22.3. Contains a 60S Ribosomal Protein L21 pseudogene and an HNRNP A3 (Heterogenous Nuclear Riboprotein A3, FBRNP) pseudogene. Contains ESTs, STSs and GSSs, complete sequence //13.3e-05.124/75//AL024509
  - F-HEMBA1000752//Human Chromosome X, complete sequence.//5.9e-48:502:75//AC004073
    - F-HEMBA1000769//Homo sapiens clone NH0576N21, WORKING DRAFT SEQUENCE, 5 unordered pieces.//
      0.011:179:67//AC005043
- 35 F-HEMBA1000773//Caenorhabditis elegans DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone Y59A8, WORKING DRAFT SEQUENCE.//0.070:231:63//Z98870
  - F-HEMBA1000774//Homo sapiens PAC clone DJ1059M17 from 7q21-q31.1, complete sequence.//6.2e-40:385: 75//AC004953
    - F-HEMBA1000791
- 4º F-HEMBA1000817//Myrmecia pilosula HI87-135 mitochondrion cytochrome b gene, partial cds.//0.99:244:58// LI15678
  - F-HEMBA1000822//Human DNA sequence from PAC 179D3, between markers DXS6791 and DXS8038 on chromosome X contains S10 GTP-binding protein, ESTs and CpG island.//0.033:294:62//Z81370
- F-HEMBA1000827/lborrella burgdorferi (section 50 of 70) of the complete genome./l/3-Fe-05463.58/l/AE001164
  F-HEMBA1000843//Homo sapiens DNA sequence from clone 511824 on chromosome 20q11.2-12. Contains the TOP1 gene for Topoisomerase I, the PLCG1 gene for 1-Phosphatidylinositio-4,5-Bisphosphate Phosphodiesterase Gamma 1 (EC 3.1.4.11, PLC-Gamma-1, Phospholipase C-Gamma-1 PLC-II, PLC-148), the KIAA0395 gene for a probable Zinc Finger Homeobox protein and a 60S Ribosomal Protein L23 LIKE pseudogene. Contains a predicted CpG island, ESTs, STSs and GSSs, complete sequence./l/3.0e-155:732-88/IAL022394
- F-HEMBA1000851//Rattus norvegicus glucocorticoid modulatory element binding protein 2 mRNA, complete cds.// 1.6e-31:386:72//AF059273
  - F-HEMBA1000852//Homo sapiens Xp22 bins 3-5 PAC RPCI4-617A9 (Roswell Park Cancer Institute Human PAC Library) containing Arylsulfatase D and E genes, complete sequence.//8.5e-115.455:98//AC005295 F-LHEMBA100887
- 55 F-HEMBA1000868//Human DNA sequence from cosmid J138O17, between markers DXS6791 and DXS8038 on chromosome X contains EST CA repeat and an endogenous retroviral like element.//i6.6e-41424.75//Z72519 F-HEMBA1000870//Gnamptodon pumilio cytochrome oxidase II gene, partial ods; and tRNA-Asp, tRNA-His, and tRNA-Lys genes, complete sequence, mitochondrial genes for mitochondrial products.//i0.0049:21166//AF034598

- F-HEMBA1000872//CIT-HSP-2355D20.TF CIT-HSP Homo sapiens genomic clone 2355D20, genomic survey sequence //3 7e-33:180:98//AQ059583
- F-HEMBA1000876//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 473B4, WORKING DRAFT SEQUENCE.//5.6e-37:262:72//Z83826
- F-HEMBA1000908//Triticum aestivum low-affinity cation transporter (LCT1) mRNA, complete cds.//1.0:304:59//
  - AF015523
    F-HEMBA1000910//M.musculus necdin mRNA\_complete cds.//6.1e-08:256:61//M80840
  - F-HEMBA1000918//Tetrahymena thermophila micronuclear developmentally eliminated sequence region.//0.13: 232:63//188158
- 10 F-HEMBA1000919//Gallus domesticus filamin mRNA, complete cds.//1.0:213:65//U00147
  - F-HEMBA1000934//CIT-HSP-2053H24.TR CIT-HSP Homo sapiens genomic clone 2053H24, genomic survey sequence.//5.5e-11:275:64//B69224
  - F-HEMBA1000942//Homo sapiens clone DJ0754G14, WORKING DRAFT SEQUENCE, 15 unordered pieces.// 9.7e-05:78:83//AC004878
- 15 F-HEMBA1000943//Homo sapiens chromosome 17, clone hRPK.640\_I\_15, complete sequence.//5.8e-140:661: 99//AC005324
  - F-HEMBA1000946
    - F-HEMBA1000960//Homo sapiens clone DJ1111F22, WORKING DRAFT SEQUENCE, 12 unordered pieces.//
      8.3e-16:181:75//AC004967
- F-HEMBA100968//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 69M21, WORKING DRAFT SEQUENCE //4.4e-117:398:86//AL031735
  - F-HEMBA1000971//H.sapiens CpG island DNA genomic Mse1 fragment, clone 182f4, forward read cpg182f4 ff1a.//1.5e-20:126:96//Z57528
- F-HEMBA1000972//Homo sapiens genomic DNA of 8p21.3-p22 anti-oncogene of hepatocellular colorectal and non-small cell lung cancer, segment 1/11.//0.34;642:59//AB020858
  - F-HEMBA1000974//Homo sapiens clone DA0091H08, complete sequence.//5.1e-183:865:98//AC004817
    - F-HEMBA1000975//Orf virus homologue of retroviral pseudoprotease gene, complete cds.//0.00065:391:62// M30023
- F-HEMBA1000885 I/Human DNA sequence from clone 272E8 on chromosome Xp22.13-22.31. Contains a pseudogene similar to MDM2-Like P53-binding protein gene. Contains STSs, GSSs and a CA repeat polymorphism, complete sequence (I/3 4e-05:24766/I/293929
  - F-HEMBA1000986//Homo sapiens DNA from chromosome 19-cosmid R31491, genomic sequence.//6.6e-06:508: 61//AD000813
  - F-HEMBA1000991//Homo sapiens mRNA for Hrs, complete cds.//1.2e-22:193:84//D84064 F-HEMBA1001007
- 35 F-HEMBA1001008//Human DNA sequence from clone 391022 on chromosome 6p21.2-21.31 Contains pseudogenes similar to ribosomal protein, ESTs, GSSs, complete sequence.//7.8e-46:532:73//AL031577
  - F-HEMBA1001009//Human mRNA for IgM heavy chain complete sequence.//0.97:369:59//X17115
  - F-HEMBA1001017//Homo sapiens mRNA for KIAA0468 protein, complete cds.//4.4e-139:661:98//AB007937
    F-HEMBA1001019//Homo sapiens, clone hRPK.15\_A\_1, complete sequence,//1.6e-16:521:64//AC006213
- 40 F-HEMBA1001020//Homo sapiens chromosome 17, clone hRPK.178\_C\_3, complete sequence.//3.8e-50:367:72// AC005702
  - F-HEMBA1001022
  - F-HEMBA1001024//Homo sapiens T-cell receptor alpha delta locus from bases 1 to 250529 (section 1 of 5) of the Complete Nucleotide Sequence.//5,0e-23:378:69//AE000658
- 45 F-HEMBA1001026//Homo sapiens DNA sequence from PAC 435D1 on chromosome Xq25. Contains ESTs and STS.//7.6e-19:867:60//Z86064
  - F-HEMBA1001043//HS\_2219\_B1\_A10\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2219 Col=19 Row=B, genomic survey sequence.//3.0e-15:124:88//AQ301521
- F-HEMBA1001051/iHuman Chromosome X clone bWXD342, complete sequence.//4.8e-79:308.84/iAC004072 F-HEMBA1001052/iHomo sapiens chromosome 17, clone hRPK.146\_P\_2, WORKING DRAFT SEQUENCE, 4 unordered pieces.// 05.3384:61/iAC005341
  - F-HEMBA1001059//Human N-acetylgalactosamine 6-sulphatase (GALNS) gene, exon 10.//2.8e-26:397:71//
- F-HEMBA1001060//Homo sapiens chromosome 17, clone hRPK.855\_D\_21 complete sequence.//0.98:280:62//
  AC006079
- F-HEMBA1001071//Human mRNA for pro alpha 1 (III) collagen C-terminal propeptide.//1.1e-31:181:96//X01742
  - F-HEMBA1001077//nuclear protein TIF1 [mice, mRNA, 3951 nt].//3.6e-13:338:65//S78219
  - F-HEMBA1001080//Streptomyces coelicolor cosmid 1A9.//0.00012:364:63//AL034446

- F-HEMBA1001085//Human Chromosome 15q26.1 PAC clone pDJ290i21 containing fur, fes, and alpha mannosidase IIx genes, WORKING DRAFT SEQUENCE, 9 unordered pieces. I/8.5e-134:476:96//AC004586
- F-HEMBA1001088//Sequence 1 from patent US 5552529.//2.2e-71:303:78//I25863
- F-HEMBA1001094//Homo sapiens clone RG491N20, complete sequence.//8.9e-119:609:96//AC005105
- F-HEMBA1001099
  - F-HEMBA1001109//Homo sapiens BAC clone RG318M05 from 7g22-g31.1, complete sequence.//2.4e-58:347: 87//AC005250
  - F-HEMBA1001121//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 90G24, WORKING DRAFT SEQUENCE.//3.4e-21:226:65//AL008723
- 10 F-HEMBA1001122//Plasmodium falciparum chromosome 2, section 20 of 73 of the complete sequence.//9.2e-07: 732:57//AF001383
  - F-HEMBA1001123//Homo sapiens full-length insert cDNA clone ZD38E12.//1.1e-11:231:68//AF086247
  - F-HEMBA1001133//Homo sapiens clone DJ0856O24, WORKING DRAFT SEQUENCE, 4 unordered pieces.// 0.011:163:69//AC004909
- 15 F-HEMBA1001137//Homo sapiens mRNA for KIAA0798 protein, complete cds.//6.9e-72:527:77//AB018341
  - F-HEMBA1001140//Homo sapiens clone RG228D17, WORKING DRAFT SEQUENCE, 2 unordered pieces. I/2. 3e-
    - F-HEMBA1001172//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-628 genomic seguence, WORKING DRAFT SEQUENCE, 9 unordered pieces, I/O, 010:520:59//AC005507
- 20 F-HEMBA1001174//R.norvegicus (Sprague Dawley) ARL5 mRNA for ARF-like protein 5.//1.0e-59:565:73//X78604 F-HEMBA1001197//Homo sapiens clone 82F9, WORKING DRAFT SEQUENCE, 4 unordered pieces, //0.0037; 151:70//AC004815
  - F-HEMBA1001208//Human BAC clone RG264L19 from 7p15-p21, complete sequence.//7.4e-35:195:81// AC002410
- 25 F-HEMBA1001213//Homo sapiens clone DJ0892G19, complete seguence, //1.9e-171:826:98//AC004917
  - F-HEMBA1001226//Homo sapiens clone DJ0850101, WORKING DRAFT SEQUENCE, 1 unordered pieces.// 0.00010:557:57//AC006009
  - F-HEMBA1001235//Homo sapiens chromosome 17, clone hRPK.601 N 13, complete sequence.//0.0086:372: 58//AC005389
- 30 F-HEMBA1001247//H.sapiens CpG island DNA genomic Mse1 fragment, clone 11b11, reverse read cpg11b11.rt1a.//2.0e-24:154:93//Z64441
  - F-HEMBA1001257//Homo sapiens alpha-methylacyl-CoA racemase mRNA, complete cds.//1.9e-88:659:81// AF047020
    - F-HEMBA1001265//Human 18S ribosomal RNA.//1.0e-32:180:97//X03205
- 25 F-HFMBA1001281

- F-HEMBA1001286//B.taurus mRNA for RF-36-DNA-binding protein,//7.7e-26:236:81//X15543
- F-HEMBA1001289//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-69G12, complete sequence.//5.5e-28:530:64//AC004131
- F-HEMBA1001294//Yeast mitochondrial appligene for ATPase subunit 8.//2.8e-15:722:60//X00960
- 40 F-HEMBA1001299//Human DNA sequence from clone 422G23 on chromosome 6q24 Contains EST, STS, GSS, CpG island, complete sequence.//4.2e-24:288:76//AL031003
  - F-HEMBA1001302/cDNA encoding a human homologue of a mouse novel polypeptide derived from stromal cell.// 7.2e-121:439:96//E12260
- F-HEMBA1001303//Plasmodium falciparum 3D7 chromosome 12 PFYAC181 genomic sequence, WORKING 45 DRAFT SEQUENCE, 8 unordered pieces, 1/0, 011:637:56//AC005505
  - F-HEMBA1001310//HS 3252 B2 B12 T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3252 Col=24 Row=D, genomic survey sequence.//1.2e-16:166:82//AQ217054
    - F-HEMBA1001319//CIT-HSP-2034J6.TF CIT-HSP Homo sapiens genomic clone 2034J6, genomic survey sequence.//0.33:256:59//B79408
- 50 F-HEMBA1001323//Homo sapiens proto-oncogene (Wnt-5a) mRNA, complete cds.//7.8e-30:165:99//L20861 F-HEMBA1001326//Homo sapiens DNA sequence from BAC 55C20 on chromosome 6. Contains a Spinal Muscular Atrophy (SMA3) LIKE gene overlapping with a beta-glucoronidase LIKE pseudogene. Contains a membrane protein LIKE pseudogene, a Glyceraldehyde 3-phosphate dehydrogenase (GAPDH) LIKE pseudogene, five predicted tRNA genes, Contains ESTs, GSSs(BAC end sequences) and a CA repeat polymorphism, complete se-
- quence.//5.4e-19:347:68//AL021368 F-HEMBA1001327//CIT-HSP-2354E10.TR CIT-HSP Homo sapiens genomic clone 2354E10, genomic survey sequence.//0.012:152:65//AQ075713
  - F-HEMBA1001330//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from contig 3-103, com-

plete sequence //0.0037:254:62//AL010208

F-HEMBA1001351//Homo sapiens VAMP-associated protein of 33 kDa (VAP-33) mRNA, complete cds.//1.1e-103: 516:97//AF057358

F-HEMBA1001361//Homo sapiens chromosome 9, clone hRPK.202\_H\_3, complete sequence.//1.7e-150:706:99// AC006241

F-HEMBA1001375//Streptomyces coelicolor cosmid 1E6.//1.0:375:59//AL033505

F-HEMBA1001377//HS\_3020\_B1\_D12\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3020 Col=23 Row=H, genomic survey sequence://0.00022:63:77//AQ105297

F-HEMBA1001383//Plasmodium falciparum chromosome 2, section 68 of 73 of the complete sequence.//0.00035: 317:60//AE001431

517:60/AE001431 F-HEMBA1001387/HS\_3039\_B1\_D01\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3039 Col=1 Row=H, genomic survey sequence://5.0e-90.437:88//AO155035

F-HEMBA1001388/Homo sapiens done RG189J21, WORKING DRAFT SEQUENCE, 15 unordered pieces.//
4 2e.47 156/89/IAC069073

F-HEMBA1001391//Human DNA sequence from clone 409O10 on chromosome 20q12 Contains CA repeat, GSS, STS, complete sequence.//2.0e-06:495:60//AL031256

F-HEMBA1001398//H.sapiens CpG island DNA genomic Mse1 fragment, clone 70d11, forward read cpg70d11.ft1b.//0.018:46:97//Z62591

F-HEMBA1001405//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 50024, WORKING DRAFT SEQUENCE //2 3e-74-623:71//AI 034380

F-HEMBA1001407//Mus musculus domesticus Torino (Sry) gene, complete cds.//0.36:363:57//U03645

F-HEMBA1001411//Homo sapiens genomic DNA, 21q region, clone: S39BG29, genomic survey sequence.//8.4e-12:516:60//AG001050

F-HEMBA1001413

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25 F-HEMBA1001415//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 41018, WORKING DRAFT SEQUENCE.//0.98:177:64//AL031732

 $\label{eq:first-problem} F-HEMBA1001432//Homo sapiens clone DJ0693M11, WORKING DRAFT SEQUENCE, 7 unordered pieces.// \\ 8.0e-177:859:97//AC006146$ 

F-HEMBA1001433//Homo sapiens clone DJ0892G19, complete sequence.//2.0e-35:376:64//AC004917

30 F-HEMBA1001435//Homo sapiens chromosome 17, clone hRPK.63\_A\_1, complete sequence.//1.2e-74:284:84// AC005670

F-HEMBA1001442//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from contig 3-66, complete sequence.//0.056:194:63//AL010138

F-HEMBA1001446//Homo sapiens chromosome 4 clone B150J4 map 4q25, complete sequence.//0.96:328:61// AC004047

F-HEMBA1001450

F-HEMBA1001454//Human DNA sequence from clone 598A24 on chromosome Xp11.1-11.23 Contains zinc finger X-linked proteins ZXDA, ZXDB, ESTs and STS, complete sequence.//2.0e-47:468:73//AL031115

F-HEMBA1001455//CIT978SK-32J2.TV CIT978SK Homo sapiens genomic clone 32J2, genomic survey sequence.//1.5e-05:223:65//B78859

F-HEMBA1001463//cSRL-69d1-u cSRL flow sorted Chromosome 11 specific cosmid Homo sapiens genomic clone cSRL-69d1, genomic survey sequence.//5.1e-66:564:77//B05652

F-HEMBA1001476//Homo sapiens mRNA for KIAA0572 protein, partial cds.//1.9e-102:489:99//AB011144

F-HEMBA1001478/HS\_2228\_A2\_B03\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2228\_Cole6 Row=C, genomic survey sequence./I4.5e-40.275.88/IAQ032041 F-HEMBA1001497/Human DNA sequence from clone 281H8 on chromosome 6q25.1-25.3. Contains up to four

F-HEMBA1001497/Human DNA sequence from clone 281H8 on chromosome 6q25.1-25.3. Contains up to four novel genes, one with similarity to KIAA0323 and worm C39712.1 and another with Ubliquith-Like protein gene SMT3 (the latter in an intron of a novel gene). Contains ESTs, STSs, GSSs, a putative CpG island and genomic marker D6S1553, complete sequence //T.7-64-7311:86/MA.031133

F-HEMBA1001510//Human HLA class III region containing cAMP response element binding protein-related protein (CREB-RP) and tenascin X (tenascin-X) genes, complete cds, complete sequence.//2.0e-130:699.93//U89337 F-HEMBA1001515//Homo sapiens chromosome 19, cosmid F24866, complete sequence.//4.1e-114:711:85// AC005794

F-HEMBA1001517//Homo sapiens BAC clone RG459N13 from 7p15, complete sequence.//5.7e-162:769:98// AC004549

F-HEMBA1001522//Caenorhabditis elegans cosmid ZK328.//8.6e-17:498:61//U50193

F-HEMBA1001526//Human DNA sequence from cosmid 444G9 from a contig from the tip of the short arm of chromosome 16, spanning 2Mb of 16p13.3 Contains ESTs and CpG islands,.//0.31:120:69//Z98258

- F-HEMBA1001533
- F-HEMBA1001557//Chionoecetes opilio (clone COP41) DNA microsatellite repeat regions.//7.0e-25:303:72//
- F-HEMBA1001566//Homo sapiens DNA sequence from PAC 12703 on chromosome 1q23-25. Contains FMO2 and FMO3 genes for Flavin-containing Monooxygenase 2 and Flavin-containing Monooxygenase (N-Oxide 3, EC1.14.138, Dimethylaniline Oxidase 3, FMO II, FMO 3), and a gene for another, unknown, Flavin-containing Monooxygenase family protein. Contains ESTs and GSSs, complete sequence/T/2-e148056-06/ML021026
- F-HEMBA1001569//Homo sapiens mRNA for vesicle associated membrane protein 2 (VAMP2).//1.1e-64:338:95// AJ225044
  - F-HEMBA1001570//Homo sapiens PAC clone DJ0844F09 from 7p12-p13, complete sequence.//2.1e-148:698:
  - F-HEMBA1001579//Homo sapiens mRNA for NS1-binding protein (NS1-BP).//2.2e-173:678:99//AJ012449
  - F-HEMBA1001581//Homo sapiens clone DJ1158B01, WORKING DRAFT SEQUENCE, 23 unordered pieces.// 0.30:484:59//AC004980
- F-HEMBA1001585

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- F-HEMBA1001589//Human BAC clone RG317G18 from 7q31, complete sequence.//0.98:197:63//AC002432
- F-HEMBA1001595//Human mRNA for KIAA0128 gene, partial cds.//8.2e-109:855:78//D50918
- F-HEMBA1001608//RPCI11-72E2.TJ RPCI11 Homo sapiens genomic done R-72E2, genomic survey sequence.// 3.8e-05:235:64//AQ267131
- F-HEMBA1001620//Oryza sativa RINO1 mRNA for myo-inositol phosphate synthase, complete cds.//3.8e-40:719: 64//AB012107
- F-HEMBA1001635//HS\_3208\_A1\_D07\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3208 Col=13 Row=G, genomic survey sequence.//1.4e-15:120:90//AQ176944
- F-HEMBA1001636//Homo sapiens 12q24 PAC RPCI1-66E7 (Roswell Park Cancer Institute Human PAC library) complete sequence.i/0.15:221:64//AC004216 F-HEMBA1001640/HS 3258 B2 DO3 MR CIT Approved Human Genomic Sperm Library D Homo sapiens generated the page 12 point of the page 12 point
  - nomic clone Plate=3253 Co1=6 Row=H, genomic survey sequence //9.1e-52:278:95//AQ216058
    F-HEMBA1001647//H. sapiens gene for plectin.//0.00052:629:61//254367
- 30 F-HEMBA1001651//Salmo salar DNA for a cryptic repeat.//7.9e-08:270:64//AJ012206
- F-HEMBA1001655//Homo sapiens chromosome 5, BAC clone 194j18 (LBNL H158), complete sequence.//5.9e-
  - F-HEMBA1001655/iHomo sapiens chromosome 5, BAC clone 194J18 (LBNL H158), complete sequence.//5.9e 164:802:97//AC005368
    - F-HEMBA1001658//M.musculus COL3A1 gene for collagen alpha-I.//2.4e-30:742:62//X52046
  - - F-HEMBA1001672//Homo sapiens methyl-CpG binding protein MBD3 (MBD3) mRNA, complete cds.//6.1e-152 725:98//AF072247
  - F-HEMBA1001675//RPCI11-54F8.TV RPCI11 Homo sapiens genomic clone R-54F8, genomic survey sequence.//
    5.3e-75:341:85//AQ082126
- 40 F-HEMBA1001678//Homo sapiens Xp22 PAC RPCI1-167A22 (from Roswell Park Cancer Center) complete sequence.//8.4e-54:551:74//AC002349 F-IF-MRA1001681
  - F-HEMBA1001702//Plasmodium falciparum chromosome 2, section 35 of 73 of the complete sequence.//0.94: 676:54//AF001398
- 45 F-HEMBA1001709//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 702J19, WORKING DRAFT SEQUENCE //0.74:659:58//AL033531
  - F-HEMBA1001711//Lysiphlebus melandriicola NADH dehydrogenase 1 gene, mitochondrial gene encoding mitochondrial protein, partial cds.//3.0e-07:413:60//AF069178
    - F-HEMBA1001712//Homo sapiens BAC clone RG041H04 from 7q21-q22, complete sequence.//0.091:315:61// AC004519
    - F-HEMBA1001714//Rattus norvegicus mitochondrial ATPase inhibitor gene, complete cds.//1.6e-28:218:75//
      - F-HEMBA1001718//HS\_3056\_A2\_H08\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3056 Col=16 Row=O, genomic survey sequence.//2.0e-79:383:99//AQ106367
- F-HEMBA1001723//HS\_2188\_A2\_D02\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2188 Col=8 Row=G, genomic survey sequence.//3.8e-28:174:94//AQ116793
  - F-HEMBA1001731//HS\_3021\_A1\_A11\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3021 Col=21 Row=A, genomic survey sequence.//2.5e-11:420:62//AQ154658

- F-HEMBA1001734//Homo sapiens chromosome Y, clone 264,M,20, complete sequence.//0.00060:392:60// AC004617
- F-HEMBA1001744//HS\_3194\_A1\_D05\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3194 Col=9 Row=G, genomic survey sequence.//5.8e-29:163:97//AQ252295
- F-HEMBA1001745//Homo sapiens chromosome 9q34, clone 280C11, complete sequence.//0.66:627:59// AC002102
  - F-HEMBA1001746//HS\_2163\_B1\_F04\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2163 Col=7 Row=L, genomic survey seguence.//1.4e-16:238:70//AQ085995
  - F-HEMBA1001761//Genomic sequence from Mouse 9, complete sequence.//3.5e-52:198:86//AC002109
- 10 F-HEMBA1001781
  - F-HEMBA1001784//Genomic sequence from Human 9q34, WORKING DRAFT SEQUENCE, 2 unordered-pieces.//5.5e-13:296:65//AC002099
  - F-HEMBA1001791//Homo sapiens DNA from chromosome 19-cosmids R31158, R31874, and R28125, genomic sequence, complete sequence.//0.18:534:59//AF038458
- F-HEMBA1001800//CrT-HFP-2049N5.TF CIT-HSP Homo sapiens genomic clone 2049N5, genomic survey sequence.//2.2e-40:335:80//AQ009222
  - F-HEMBA1001803//M.musculus (Ba1b/C) P/L01 mRNA.//1.7e-25:286:74//Z31360
    - F-HEMBA1001804//Mouse interleukin 2 receptor (p55 IL-2R) mRNA, 5' end.//1.9e-58:358:89//M21977
  - F-HEMBA1001808//Homo sapiens mRNA, chromosome 1 specific transcript KIAA0500.//7.8e-174:809:98// AB007969
    - F-HEMBA1001809//Bovine herpesvirus 1 complete genome.//9.0e-09:639:57//AJ004801
    - F-HEMBA1001815

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- F-HEMBA1001819//HS\_3079\_B1\_E04\_T7 CIT Approved Human Genomic Sperm Library D Homo saplens genomic clone Plate=3079 Col=7 Row=J, genomic survey sequence.//1.4e-79:396:97//AQ186616
- 25 F-HEMBA1001820//Homo sapiens BAC clone GS165L15 from 7p15, complete sequence.//0.00026:436:60// AC005013
  - F-HEMBA1001822//Homo sapiens intersectin short form mRNA, complete cds.//1.2e-40:510:65//AF064243
  - F-HEMBA1001824//Homo sapiens expanded SCA7 CAG repeat.//6.1e-20:344:68//AF020275
    F-HEMBA1001835//Homo sapiens BAC clone RG017K18 from 7g31, complete sequence.//0.0094:553:58//
- 30 AC005161
  - F-HEMBA1001844//Homo sapiens chromosome Xp22-135-136 clone GSHB-56711, WORKING DRAFT SE-QUENCE, 35 unordered pieces.//1.2e-22:316:70//AC005867
    - F-HEMBA1001847//M.musculus Zfp-29 gene for zinc finger protein.//5.3e-27:397:69//X55126
    - F-HEMBA1001861//Homo sapiens mRNA for KIAA0617 protein, complete cds.//8.8e-184:865:98//AB014517
- 35 F-HEMBA1001864//Arabidopsis thaliana chromosome II BAC F17H15 genomic sequence, complete sequence.// 0.38:337:62//AC005395
  - F-HEMBA1001866//Caenorhabditis elegans cosmid F48E3.//1.4e-10:224:63//U28735
  - F-HEMBA1001869//Homo sapiens BA $\bar{\rm C}$  clone RG114B19 from 7q31.1, complete sequence.//6.7e-98:288:91// AC005065
- 40 F-HEMBA1001888//Human Chromosome 11p15.5 PAC clone pDJ915f1 containing KvLQT1 gene, complete sequence.//4.9e-114:476:84//AC003693
  - F-HEMBA1001896//Bos taurus pyruvate dehydrogenase phosphatase regulatory subunit precursor, mRNA, complete cds.//2.2e-137:839:86//AF026954
- F-HEMBA1001910//Homo sapiens Chromosome 2p13 BAC Clone h173, complete sequence.//0.90:221:63//
  AC003065
  - F-HEMBA1001912//HS\_2237\_A1\_C10\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2237 Col=19 Row=E, genomic survey sequence.//9.7e-76:364:100//AQ033732
    - F-HEMBA1001913//Leishmania major chromosome 3 clone L4625 strain Friedlin, WORKING DRAFT SE-QUENCE, 6 unordered pieces //0.00063;219:65//AC005766
    - F-HEMBA1001915//Homo sapiens genomic DNA of 9q32 anti-oncogene of flat epitherium cancer, segment 5/10.// 0.00011:366:63//AB020873
      - F-HEMBA1001918//Pneumocystis carinii gene for major surface glycoprotein MSG105, exon1-2, complete cds.// 0.00024:562:56//D82031
- F-HEMBA1001921//Homo sapiens germinal center kinase related protein kinase mRNA, complete cds.//2.1e-184: 855:99//AF000145
  - F-HEMBA1001939/Human DNA sequence from clone 395P12 on chromosome 1q24-25. Contains the TXGP1 gene for tax-transcriptionally activated glycoprotein 1 (34kD) (0X40 ligand, 0X40L) and a GOT2 (Aspartate Autionotransferase, mitochondrial precursor, EC 2.6.1.1, Transaminase A, Glutamate Oxaloacetate Transaminase-2)

- pseudogene. Contains ESTs, STSs and GSSs, complete sequence.//1.1e-42:380:80//AL022310
  F-HEMBA1001940/Homo sapiens clone DJ1093116, WORKING DRAFT SEQUENCE, 5 unordered pieces.//7.5e-175:881:97//AC005629
- F-HEMBA1001942//Homo sapiens chromosome 12p13.3 clone RPCI1-96H9, WORKING DRAFT SEQUENCE, 66 unordered pieces.//0.097:107:71//AC006057
  - F-HEMBA1001945//Drosophila F family transposable element F12 3' region.//0.94:140:65//X01934 F-HEMBA1001950//H.sapiens CpG island DNA genomic Mse1 fragment, clone 15b5, forward read cpg15b5.ft1q.//
    - F-HEMBA1001950//H.sapiens CpG island DNA genomic Mse1 fragment, clone 15b5, forward read cpg15b5.ft1q.// 1.4e-27:168:95//Z54728
  - F-HEMBA1001960//Locusta migratoria mRNA for nAChR alpha1 subunit.//0.010:108:71//AJ000390
- F-HEMBA1001962//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-628 genomic sequence, WORKING DRAFT SEQUENCE, 9 unordered pieces.//9.7e-05:494:60//AC005507
  - F-HEMBA1001964

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- F-HEMBA1001967//Human DNA sequence from done 341E18 on chromosome 6p11.2-12.3. Contains a Serine/ Threonine Protein Kinase gene (presumptive isolog of a Rat gene) and a novel atternatively spliced gene. Contains a putative CoG island. ESTs and GSSs. complete sequence //9 6e-122:373-99//AL031178
- F-HEMBA1001979/HS\_3067\_B1\_A06\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3067 Col=11 Row=B, genomic survey sequence.//0.43:193:64//AQ143506
  - F-HEMBA1001987//Plasmodium falciparum MAL3P6, complete sequence.//1.0:428:56//Z98551
  - F-HEMBA1001991//HS\_2237\_A2\_G09\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2237 Col=18 Row=M, genomic survey sequence.//4.3e-05:240:64//AQ067283
- F-HEMBA1002003//protein phosphatase 2C isoform [rats, liver, mRNA, 1950 nt].//2.7e-33:364:74//S90449 F-HEMBA1002008//WORKING DRAFT SEQUENCE, 3 unordered pieces.//0.00032:214:88//AC005948 F-HEMBA1002018
  - F-HEMBA10020227/Human p37NB mRNA, complete cds.//0.014;58;96//U32907
- 25 F-HEMBA1002035//Mouse transcriptional control element.//7.8e-07:200:69//M17284
  - F-HEMBA1002039//Human DNA sequence from cince 267M20 on chromosome Xq22.2-22.3. Contains part of the DIAPH2 gene and a pseudogene, ESTs, STSs and GSSs, complete sequence.//ii.3149/756/ML031053 F-HEMBA1002049//Homo sapiens chromosome 5, BAC clone 28287 (LBNL H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 5, BAC clone 28287 (LBNL H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Homo sapiens chromosome 6, BAC clone 28287 (LBNL) H192), complete sequence.//ii.55e-145MBA1002049//Hippacoccomplete sequence.//iii.55e-145MBA1002049//Hippacoccomplete sequence.//iii.55e-145MBA1002049//Hi
- 42:532:63//AC005216

  F-HEMBA1002084//Homo sapiens chromosome 19 cosmid F15386, genomic sequence, complete sequence.//
- 0.81:435:59//AF025422
  F-HEMBA1002092//Mus musculus Olf-1/EBF-like-3 transcription factor (O/E-3) mRNA, complete cds.//7.2e-130:
  - 769:87//U92703
    F-HEMBA1002100//Homo sapiens PAC clone DJ0991G20, complete sequence.//1.3e-47:124:96//AC004943
- 35 F-HEMBA1002102//Xenopus laevis mRNA for xSox7 protein, complete cds.//2.7e-13:132:71//D83649 F-HEMBA1002113//F.rubripes GSS sequence, clone 063K10bB4, genomic survey sequence.//0.029:142:66// 78BB40
  - F-HEMBA1002119/Human Chromosome 11 pac pDu1173a5, complete sequence./1.3e-14:515:62//AC000378 F-HEMBA1002125/Homo sapiens calcium-activated potassium channel (KCNN3) mRNA, complete cds.//0.98: 22261/JAF031815
  - 22.2.////N/O 2019//Caenorhabditis elegans cosmid F55C9, complete sequence.//0.0081:371:60//Z81549
    F-HEMBA1002144//Saccharomyces cerevisiae mitochondrion transfer RNA-Met (tRNA-Met) gene, oxil gene, and
    ORF 1/M 9-06-3414 fil.36888
- F-HEMBA1002150//Homo sapiens mRNA for KIAA0720 protein, partial cds.//0.00017:353:62//AB018263 F-HEMBA1002151
- F-HEMBA1002153//CITBI-E1-2519120.TR CITBI-E1 Homo sapiens genomic clone 2519I20, genomic survey sequence //8 5e-61:334:94//AQ277613
  - F-HEMBA1002160//Homo sapiens clone DJ1189D06, complete sequence.//8.5e-44:385:77//AC005232
- F-HEMBA1002161//Coturnix coturnix slow myosin heavy chain 2 (qmyhc2) mRNA, partial cds.//2.1e-59:571:74// 4F006829
- F-HEMBA1002162//Homo sapiens Xp22-150 BAC GSHB-309P15 (Genome Systems Human BAC Library) complete sequence//f/3-e5-35-988-67//AC006210
  F-HEMBA1002166//Human DNA sequence from PAC 84F12 on chromosome Xq25-Xq26.3. Contains glypican-3
- precursor (intestinal protein OCI-5) (GTR2-2), ESTs and CA repeat.//1.2e-50:319:78//AL008712

  F-HEMBA1002177//Homo sapiens BAC clone RG293F11 from 7q21-7q22, complete sequence.//2.5e-18:150:88//
  - AC000066
    F-HEMBA1002185/Homo sapiens clone DJ0292L20, WORKING DRAFT SEQUENCE, 2 unordered pieces.//
    0.00066-466:59//AC004825

- F-HEMBA1002189//Homo sapiens clone GS166C05, WORKING DRAFT SEQUENCE, 7 unordered pieces. //3.3e-23:176:77//AC005015
- F-HEMBA1002191//Homo sapiens mRNA for KIAA0689 protein, partial cds.//1.0:382:59//AB014589
- F-HEMBA1002199//Homo sapiens chromosome 4 clone B55B24 map 4q25, complete sequence.//1.8e-20:368: 66//AC005150
- F-HEMBA1002204//HS\_2055\_A1\_H09\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2055 Col=17 Row=O, genomic survey sequence.//1.2e-06:178:65//AQ235350
  - F-HEMBA1002212//S.cerevisiae chromosome IV reading frame ORF YDL101c.//0.035:345:60//Z74149
  - F-HEMBA1002215//M.musculus mRNA for testin.//4.6e-80:504:87//X78989
- F-HEMBA1002226//Homo sapiens Xp22 bins 87-93 PAC RPCI1-122K4 (Roswell Park Cancer Institute Human PAC Library) complete sequence.//5.7e-63:336:74//AC003035
  - F-HEMBA1002229//Homo sapiens BAC clone NH0539B24 from 7p15.1-p14, complete sequence.//2.6e-39:311: 81//AC006044
  - F-HEMBA1002237//Homo sapiens PAC clone DJ0696N01 from 7p21-p22, complete sequence.//1.6e-12:397:64// AC004861
- F-HEMBA1002241

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  - F-HEMBA1002253
  - F-HEMBA1002257//Homo sapiens diacylglycerol kinase iota (DGKi) mRNA, complete cds.//3.5e-151:731:97// AF061936
- F-HEMBA1002265//Human DNA sequence from cosmid N28H9 on chromosome 22q11.2-qter contains ESTs, STS and endogenous retrovirus.//1.3e-09:313:62//Z71183 F-HEMBA1002267
  - F-HEMBA1002270//Homo sapiens Xp22-150 BAC GSHB-309P15 (Genome Systems Human BAC Library) complete sequence.//0.069:495:58//AC006210
- 25 F-HEMBA1002321/Homo sepiens PAC clone DJ0991023, complete sequence.//0.019:564:58//AC004944 F-HEMBA1002328//CIT-HSP-2387N15.TE.1 CIT-HSP Homo sepiens genomic clone 2387N15, genomic survey.
  - sequence.//1.8e-71:346:99//AQ240836
    F-HEMBA1002337//Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone: MYN8, complete sequence.//
- 0.84:547:57//AB020754

  30 F-HEMBA1002341//Homo sapiens mRNA for KIAA0771 protein, partial cds.//2.4e-185:872:98//AB018314
- F-HEMBA1002348//CIT-HSP-2372K24.TR CIT-HSP Homo sapiens genomic clone 2372K24, genomic survey sequence.//9.1e-33:230:75//AQ110676
  - F-HEMBA1002349//Plasmodium falciparum histidine-rich protein II (HRP II) gene, complete cds.//9.4e-06:504: 57//U69551
- 35 F-HEMBA1002363//Homo sapiens chromosome-associated protein-E (hCAP-E) mRNA, complete cds.//7.3e-188: 872:99//AF092563
  - F-HEMBA1002381//Homo sapiens genomic DNA of 8p21.3-p22 anti-oncogene of hepatocellular colorectal and non-small cell lung cancer, segment 11/11.1/2.1e-20:262;72//AB020868
  - F-HEMBA1002389//D.discoideum spore coat 60 (sp60) gene. 5' flank.//0.010:95:73//M34546
- 40 F-HEMBA1002417//Canis familiaris ZO-3 (zo-3) mRNA, complete cds.//6.2e-120:767:85//AF023617
  - F-HEMBA1002419//HS-1047-A1-F01-MF.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 830 Col=1 Row=K, genomic survey sequence.//7.6e-06:111:76//B38165
  - F-HEMBA1002430//HS\_3137\_B2\_F10\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3137 Col=20 Row=L, genomic survey sequence.//1.6e-56:367:88//AQ148697
- 45 F-HEMBA1002439//Dictyostelium discoideum actin 8 gene, 3' UTR.//0.67:129:64//M25216
  - F-HEMBA1002458//Mus musculus REX-3 mRNA, complete cds.//1.1e-30:274:72//AF051347
  - F-HEMBA1002460//Homo sapiens clone DJ1137M13, complete sequence.//4.0e-173:822:98//AC005378 F-HEMBA1002462//Sequence 41 from patent US 5708157.//9.8e-51:519:73///80067
  - F-HEMBA1002469//Human mRNA for KIAA0122 gene, partial cds.//4.0e-108:603:92//D50912
  - F-HEMBA1002475//Streptomyces coelicolor cosmid 2H4.//0.0068:626:57//AL031514
  - F-HEMBA1002477//Homo sapiens BAC clone NH0342K06 from 2, complete sequence.//1.5e-40:349:78//
  - F-HEMBA1002486

- F-HEMBA1002495//HS\_3218\_B1\_A12\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3218 Col=23 Row=B, genomic survey sequence.//1.0:179:67//AQ181410
- F-HEMBA1002498//Homo sapiens full-length insert cDNA clone ZD76B01.//1.4e-129:619:98//AF086404
  - F-HEMBA1002503//Homo sapiens clone DJ0742P04, WORKING DRAFT SEQUENCE, 6 unordered pieces.//
    1.9e-24/306/68//AC004873

- F-HEMBA1002508//Homo sapiens chromosome 19, cosmid R33516, complete sequence.//2.9e-76:464:83// AC004799
- F-HEMBA1002513//Homo sapiens mRNA for histone deacetylase-like protein (JM21).//2.8e-157:738:98//
- F-HEMBA1002515//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 407F11, WORKING DRAFT SEQUENCE //2.6e-07:307:64//AL022329
  - F-HEMBA1002538//HS\_2185\_B2\_B04\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2185 Col=8 Row=D, genomic survey sequence.//4.7e-37:339:78//AQ298315
- F-HEMBA1002542//HS\_3197\_B2\_B10\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3197 Col=20 Row=D, genomic survey sequence.//3.2e-70:372:95//AQ188792
- F-HEMBA1002547//Homo sapiens agrin precursor mRNA, partial cds.//3.5e-137:655:98//AF016903
  - F-HEMBA1002552//Human Hep27 protein mRNA, complete cds.//8.8e-07:173:68//U31875
  - F-HEMBA1002555//\*\* SEQUENCING IN PROGRESS \*\*\* Homo sapiens chromosome 4, BAC clone C0190L06; HTGS phase 1, WORKING DRAFT SEQUENCE, 21 unordered pieces, //2, 2e-15;628;60//AC004670
- F-HEMBA1002558//Human Xp22 BAC CT-285I15 (from CalTech/Research Genetics), PAC RPCI1-27C22 (from Roswell Park Cancer Center), and Cosmid U35B5 (from Lawrence Livermore), complete sequence.//2.3e-41:353: 76//AC002366
  - F-HEMBA1002561//Homo sapiens chromosome 17, clone HRPC29G21, complete sequence.//1.1e-39:538:66// AC003687
- F-HEMBA1002569//Homo sapiens protein associated with Myc mRNA, complete cds.//1.3e-140:457:99// AF075587
  - F-HEMBA1002583//CIT-HSP-2321D3.TR CIT-HSP Homo sapiens genomic clone 2321D3, genomic survey sequence.//5.1e-79:385:99//AQ038102
- F-HEMBA1002590//Homo sapiens chromosome 17, clone hRPK.167\_N\_20, complete sequence.//1.9e-35:430: 70//AC005940
  - F-HEMBA1002592//Human genomic DNA sequence from clone 308O1 on chromosome Xp11.3-11.4. Contains EST, CA repeat. STS, GSS, CpG island //4Ae-19:303;71//Z93403
  - F-HEMBA1002609//Homo sapiens mRNA for KIAA0597 protein, partial cds.//4.4e-175:820:99//AB011169 F-HEMBA1002621//Homo sapiens PAC clone DJ0650P09 from 7q21, complete sequence.//0.14:353:58//
- 30 AC004413 F-HEMBA1002624//Homo sapiens mRNA for KIAA0808 protein, complete cds.//2.9e-187:632:97//AB018351 F-HEMBA1002628//Plasmodium falciparum 3D7 chromosome 12 PFYAC812 genomic sequence, WORKING
  - DRAFT SEQUENCE, 8 unordered pieces.//1.5e-05:792:58//AC004153 F-HEMBA1002629//Streptomyces coelicolor cosmid 1A9.//8.4e-08:576:58//AL034446
- F-HEMBA1002645//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 153G14, WORKING DRAFT SEQUENCE //5.6e-47:222:86//AL031118
  - F-HEMBA1002651//Homo sapiens PAC clone DJ0593H12 from 7p31, complete sequence.//3.8e-182:859:99// AC004839
  - F-HEMBA1002659//Z.mobilis alcohol dehydrogenase I (adhA) gene, complete cds.//0.97:144:66//M32100
- 40 F-HEMBA1002661//Homo sapiens PAC clone DJ0698G21 from 7p21-p22, complete sequence.//1.3e-116:774: 84//AC004535
  - F-HEMBA1002666

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- F-HEMBA1002678//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 1137F22, WORKING DRAFT SEQUENCE.//5.7e-156:750:98//AL034421
- 45 F-HEMBA1002679//nbxb0002cC12r CUGI Rice BAC Library Oryza sativa genomic clone nbxb0002F23r, genomic survey sequence.//4.3e-09:517:58//AQ051621
  - F-HEMBA1002688//Herpes simplex virus type 2 (strain HG52), complete genome.//8.3e-20:651:61//Z86099
    F-HEMBA1002696//Mus musculus proteasome regulator PA28 beta subunit gene, complete cds.//7.6e-62:306:
    84/#EPGPA192
  - F-HEMBA1002703//Homo sapiens mRNA for KIAA0455 protein, complete cds.//1.9e-10:327:62//AB007924 F-HEMBA1002712
    - F-HEMBA1002716//HS\_3064\_A1\_C10\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3064 Col=19 Row=E, genomic survey sequence.//8.4e-97:491:96//AQ142980
- F-HEMBA1002728//Homo sapiens chromosome 5, BAC clone 205e20 (LBNL H170), complete sequence.//6.1e55 21:217:77//AC004782
  - F-HEMBA1002730//Human platelet glycoprotein IIIa (GPIIIa) gene, exon 1.//0.57:125.67/M67481
    F-HEMBA1002742/RPCI11-38J10.TP RPCI-11 Homo sapiens genomic clone RPCI-11-39J10, genomic survey sequence //1.1e-86:41499/AQ029102

- F-HEMBA1002746//Mus musculus chromosome 19, clone CIT282B21, complete sequence.//7.1e-70:303:82// AC003694
- F-HEMBA1002748//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 41018, WORKING DRAFT SEQUENCE.//0.096:212:62//AL031732
- F-HEMBA1002750//Homo sapiens chromosome 5, PAC clone 170m10 (LBNL H89), complete sequence.//6.7e-40:23270//AC004622
  - F-HEMBA1002768//Homo sapiens mRNA for KIAA0554 protein, partial cds.//9.0e-177:834:98//AB011126
  - F-HEMBA1002770//cDNA encoding novel rat protein TIP120 which is formed of complex with TBP (TATA binding protein).//1.3e-140:840:88//E12829
- F-HEMBA1002777//F.rubripes GSS sequence, clone 189C06dB12, genomic survey sequence.//1.1e-28:263:77// AI 007965
  - F-HEMBA1002779//CIT-HSP-2333I1.TF CIT-HSP Homo sapiens genomic clone 2333I1, genomic survey sequence.//1.8e-32:180:98//AO036891
  - F-HEMBA1002780//Homo sapiens PAC clone DJ0244J05 from 5q31, complete sequence.//7.0e-06:199:67// AC004592
- F-HEMBA1002794//H.sapiens mRNA for protein kinase C mu.//0.00015:244:67//X75756
  - F-HEMBA1002801//Plasmodium falciparum MAL3P2, complete sequence.//0.0010:534:57//AL034558
  - F-HEMBA1002810//Homo sapiens formin binding protein 21 mRNA, complete cds.//1.1e-167:820:97//AF071185 F-HEMBA1002816//Homo sapiens clone NH0576N21, WORKING DRAFT SEQUENCE, 5 unordered pieces.//
- 3.1e-113:254:90//AC005043 F-HEMBA1002818//Cricetulus griseus H411 precursor (H411) mRNA, complete cds.//1.2e-122:760:86//AF046870 F-HEMBA1002826//Human DNA sequence from clone 23K20 on chromosome Xq25-26.2 Contains EST, STS,
  - GSS, complete sequence.//0.0055:235:65//AL022153
    F-HEMBA1002833//Homo sapiens chromosome 17, clone hRPC.117 B 12, complete sequence.//1.4e-170:744:
- 25 99//AC004707 F-HEMBA1002850//Ephedrus persicae NADH dehydrogenase 1 gene, mitochondrial gene encoding mitochondrial
  - protein, partial cds.//1.3e-05:334:59//AF069186
    F-HEMBA1002863//CIT-HSP-2323A16.TF CIT-HSP Homo sapiens genomic clone 2323A16, genomic survey se-
- quence.//2.9e-140:750:93//AQ028419

  F-HEMBA1002876//HS\_2270\_B1\_H03\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens ge
  - nomic clone Plate=2270 Col=5 Row=P, genomic survey sequence.//0.44:163:64//AQ164031 F-HEMBA1002886
    - F-HEMBA1002896//Homo sapiens chromosome 5, P1 clone 793C5 (LBNL H58), complete sequence..//0.00015: 277:61//AC005195
- 35 F-HEMBA1002921

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- F-HEMBA1002924//CIT-HSP-2171H4.TR CIT-HSP Homo sapiens genomic clone 2171H4, genomic survey sequence.//0.0016:175:66//B89715
- F-HEMBA1002934//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 862K6, WORKING DRAFT SEQUENCE //1.26-169:797:98//AL031681
- 40 F-HEMBA1002935//Homo sapiens mRNA for KIAA0576 protein, partial cds.//4.9e-173:803:99//AB011148
  - F-HEMBA1002937//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 702J19, WORKING DRAFT SEQUENCE.//1.2e-163:411:99//AL033531
  - F-HEMBA1002939//RPCI11-74014.TJ RPCI11 Homo saplens genomic clone R-74014, genomic survey sequence.//1.7e-41:215:99//AQ266676
- 45 F-HEMBA1002944//RPCI11-55C2.TV RPCI11 Homo sapiens genomic clone R-55C2, genomic survey sequence.// 1.7e-37:375:74//AQ082240
  - F-HEMBA1002951//Homo sapiens chromosome 19, cosmid F20887, complete sequence.//0.00074:683:58// AC005578
  - F-HEMBA1002954//RPCI11-79F7.TV RPCI11 Homo sapiens genomic clone R-79F7, genomic survey sequence.// 6.1e-24:250:78//AQ284146
  - F-HEMBA1002968//HS\_2262\_B2\_G04\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2262 Col=8 Row=N, genomic survey sequence.//0.99:270:60//AQ217059
  - F-HEMBA1002970//RPCI11-5L24.TV RPCI-11 Homo sapiens genomic clone RPCI-11-5L24, genomic survey sequence.//1.4e-10:189:71//B49289
- 55 F-HEMBA1002971//CIT-HSP-2363L16.TF CIT-HSP Homo sapiens genomic clone 2363L16, genomic survey sequence.//4.3e-21:181:80//AQ080538
  - F-HEMBA1002973//Rattus norvegicus Wistar 3',5'-cyclic AMP phosphodiesterase (PDE4-10) gene, exon 10.// 2.5e-40:257:89//U01290

- F-HEMBA1002997//CIT-HSP-2387H15.TF.1 CIT-HSP Homo sapiens genomic clone 2387H15, genomic survey sequence.//9.5e-17:128:92//AQ240797
- F-HEMBA1002999//Rattus norvegicus lamina associated polypeptide 1C (LAP1C) mRNA, complete cds.//3.1e-62:713:73//U20286
- F-HEMBA1003021//Homo sapiens clone DJ0847008, WORKING DRAFT SEQUENCE, 3 unordered pieces.//7.5e-50:331:85//AC005484
  - F-HEMBA1003033//Drosophila melanogaster, chromosome 3L, region 62A10-62B5, P1 clones DS02777, DS03222, DS02345, and DS04808, complete sequence.//2.6e-20:357:66//AC005557
- F-HEMBA1003034//Human DNA sequence from 4PTEL, Huntington's Disease Region, chromosome 4p16.3.//
  4 5e-60:415:73//Z95704
  - F-HEMBA1003035//Homo sapiens chromosome Y, clone 264,M,20, complete sequence.//2.3e-05:591:57// AC004617
  - F-HEMBA1003037//RPCI11-88F2.TJ RPCI11 Homo sapiens genomic clone R-88F2, genomic survey sequence.//
    0.68:230:60//AQ286677
- 15 F-HEMBA1003041//Homo sapiens PAC clone DJ1163J12 from 7q21.2-q31.1, complete sequence //8.1e-128:550: 94//AC004983
  - F-HEMBA1003046//Homo sapiens mitochondrial processing peptidase beta-subunit mRNA, complete cds.//1.0e-164:777:98//AF054182
- F-HEMBA1003064//Plasmodium falciparum 3D7 chromosome 12 PFYAC181 genomic sequence, WORKING DRAFT SEQUENCE, 8 unordered pieces.//1.5e-07:744:59//AC005505
- F-HEMBA1003067//Rat dynorphin gene, exon 3.//1.0:140:63//M32783
  - F-HEMBA1003071/I/Homo saplens alpha2-C4-adrenergic receptor gene, complete cds.//1.5e-20:595:65//U72648 F-HEMBA1003077/I/CIT-HSP-2366/J21.TF CIT-HSP Homo saplens genomic clone 2366/J21, genomic survey sequence.//4.4e-33:176:99/IAQ080257
- F-HEMBA1003078//Homo sapiens DNA sequence from PAC 262D12 on chromosome 1q23.3-24.3. Contains a Tenascin (Hexabrachion, Cytotactin, Neuronectin, Myotendinous antigen)-LIKE gene and a mitochondrial/chloroplast 30S ribosomal protein S14-LIKE gene preceeded by a CpG island. Contains ESTs, genomic marker D1S2691 and STSs.//9.4e-43/478/70//Z99297
- F-HEMBA1003079//Homo sapiens Xp22-132-134 BAC GSHB-590J15 (Genome Systems Human BAC library)
  complete sequence.//0.96:57:85//AC004673
- F-HEMBA1003083//Homo sapiens PAC clone DJ1182N03 from 7q11.23-q21.1, complete sequence.//8.0e-74:359: 81//AC004548
  - F-HEMBA1003086//Homo sapiens chromosome 16 BAC clone CIT987SK-334D11 complete sequence.//3.6e-11: 734:58//AF001550
  - F-HEMBA1003096//Sequence 4 from patent US 5440017.//5.7e-56:594:71//I13750
    - F-HEMBA1003098//Human DNA sequence from cosmid SRL11M20, chromosome region 11p13. Contains EST and STS.//1.9e-09:230:69//Z83308
  - F-HEMBA1003117//Mouse TIS11 primary response gene, complete cds.//0.00054:480:60//M58564
  - F-HEMBA1003129//HS\_3139\_B2\_F05\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3139 Col=10 Row=L, genomic survey sequence.//2.3e-100:510:97//AQ187635
  - F-HEMBA1003133//Mouse BAC CitbCJ7 219m7, genomic sequence, complete sequence.//1.3e-78:370:90// AC005259
    - F-HEMBA1003136

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- F-HEMBA1003142//Homo sapiens full-length insert cDNA clone ZC39B06.//6.9e-121:563:100//AF086197
- 45 F-HEMBA1003148//Homo sapiens mRNA for dachshund protein.//6.7e-183:850:99//AJ005670
  - F-HEMBA1003166//Homo sapiens Chromosome 16 BAC clone CIT987-SKA-345G4 ~complete genomic sequence, complete sequence.//3.8e-27:229:76//AC002302
  - F-HEMBA1003175//Homo sapiens genomic DNA for centromeric end of MHC class I region on chromosome 6, WORKING DRAFT SEQUENCE.//9.4e-09:837:58//AB000882
- 59 F-HEMBA1003179//Homo saplens DNA sequence from Fosmid 27C3 on chromosome 22q11.2-qter. Contains two possibly alternatively spliced unknown genes, one with homology to a worm protein. Contains ESTs, complete sequence //5.4e-115.174-98//LAL023325
  - F-HEMBA1003197//Arabidopsis thaliana chromosome II BAC F15K20 genomic sequence, complete sequence.//
    1.1e-05:473:59//AC005824
- F-HEMBA1003199//Rattus norvegicus Sprague-Dawley thyroid hormone receptor alpha gene, exon 1.//1.6e-05: 367:61//J.109302
  - F-HEMBA1003202//Homo sapiens BAC clone RG437L15 from 8q21, complete sequence.//9.0e-23:247:73// AC004003

- F-HEMBA1003204//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 409J21, WORKING DRAFT SEQUENCE.//4.7e-26:141:83//Z83824
- F-HEMBA1003212//Human Chromosome 11 Overlapping Cosmids cSRL72g7 and cSRL140b8, complete sequence.//1.9e-31:158:86//AC002037
- F-HEMBA1003220//Homo sapiens chromosome 17, clone hRPC.971\_F\_3, WORKING DRAFT SEQUENCE, 1 ordered pieces.//3.4e-24:284:75//AC004150
  - F-HEMBA1003222//RPCI11-47P17.TJ RPCI11 Homo sapiens genomic clone R-47P17, genomic survey sequence.//8.7e-39:202:99//AQ202885
- F-HEMBA1003229//Arabidopsis thaliana genomic DNA, chromosome 3, P1 clone: MEB5, complete sequence.//
  0.86:227:62//AB019230
  - F-HEMBA1003235//Plasmodium falciparum chromosome 2, section 10 of 73 of the complete sequence.//8.6e-05: 372:61//AE001373
  - F-HEMBA1003250//HS-1063-A1-H02-MR.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 796 Col=3 Row=O, genomic survey sequence.//0.00032:57:96//B46142
- 15 F-HEMBA1003257//H.sapiens mRNA for RDC-1 POU domain containing protein.//2.2e-08:531:59//X64624
  - F-HEMBA1003273/H, saplens flow-sorted chromosome 6 HindIII/fragment, SC6pA19H4/I/0.070:267:64I/Z78949
    F-HEMBA1003276//CIT-HSP-2301B4.TF CIT-HSP Homo sapiens genomic clone 2301B4, genomic survey sequence./l/5.2e-08:295.63/I/AQ015073
- F-HEMBA1003278//HS\_3075\_A1\_G09\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3075 Col=17 Row=M, genomic survey sequence.//0.98:399:58//AQ120599
  - F-HEMBA1003281//High throughput sequencing of human chromosome 12, WORKING DRAFT SEQUENCE, 1 ordered pieces.//4.8e-101:277:97//AC005840
    - F-HEMBA1003286//Homo sapiens chromosome 3q13 beta-1,4-galactosyltransferase mRNA, complete cds.// 9.0e-145:539:97//AF038662
- F-HEMBA1003291//Homo sapiens mRNA for KIAA0537 protein, complete cds://5.0e-166:799:98//AB011109 F-HEMBA1003296//CITBE-E1-2507MB.TR CITBI-E1 Homo sapiens genomic clone 2507M8, genomic survey sequence://1.9e-05:388-63//AD262551
  - F-HEMBA1003304//Budworm mitochondrial partial transfer RNA-Met (tRNA-Met) gene, and partial 12S ribosomal RNA (12S rRNA) gene.//8.0e-05:388:62//L17343
- 30 F-HEMBA1003309//Crassostrea gigas clone CN20 microsatellite sequence.//0.0017:210:64//AF051177
  - F-HEMBA1003314//Homo sapiens mRNA for leucine zipper bearing kinase, complete cds.//4.6e-188:865:99// AB001872
    - F-HEMBA1003322//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 16915, WORKING DRAFT SEQUENCE.//2.4e-54:316:87//Z93015
- F-HEMBA1003327//CIT-HSP-2024C24.TRB CIT-HSP Homo sapiens genomic clone 2024C24, genomic survey sequence.//8.4e-12:166:76//B67147
  - F-HEMBA1003328//HS\_2230\_B2\_H08\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2230 Col=16 Row=P, genomic survey sequence.//0.026:128:71//AQ153313
- F-HEMBA1003330//Homo sapiens wbscr1 (WBSCR1) and replication factor C subunit 2 (RFC2) genes, complete cds.//4.0e-160:745:99//AF045555
  - F-HEMBA1003348/HS\_3194\_A1\_G05\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3194 Col=9 Row=M, genomic survey sequence.//5.0e-79:381:99//AQ173779 F-HEMBA1003369/H.vulgare GAA-satellite DNA\_//0.12:89:71/I/Z50100
    - F-HEMBA1003370//Homo sapiens cosmid 123E15, complete sequence.//3.5e-32:199:80//AF024533
- 45 F-HEMBA1003373//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 537K23, WORKING DRAFT SEQUENCE //0.019:117:71//AL034405
  - F-HEMBA1003376//Human clone HS4.66 Alu-Ya5 sequence.//4.2e-30:196:85//U67229
    - F-HEMBA1003380//Homo sapiens DNA sequence from clone 394P21 on chromosome 1p36.12-36.13. Contains the PAX7 gene, locus D1S2644, ESTs and STSs, complete sequence.//4.6e-22:206:81//AL021528
- F-HEMBA1003384//Homo sapiens clone GS096J14, WORKING DRAFT SEQUENCE, 3 unordered pieces.// 0.00094;72:90//AC006026
  - F-HEMBA1003395//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P1, WORKING DRAFT SEQUENCE.//0.00041:826:57//AL031744
- F-HEMBA1003402//CIT-HSP-2339K16.TR CIT-HSP Homo sapiens genomic clone 2339K16, genomic survey seguence.//2.4e-05:265:64//AQ056234
  - F-HEMBA1003403//Homo sapiens chromosome 4 clone B353C18 map 4q25, complete sequence.//4.3e-135:780: 98//AC004066
    - F-HEMBA1003408

- F-HEMBA1003417//Human DNA sequence from clone 496N17 on chromosome 6p11.2-12.3 Contains EST, GSS, complete sequence.//1.9e-41:239:95//AL031321
- F-HEMBA1003418//Rattus norvegicus Wistar polymeric immunoglobulin receptor (PIGR) gene, 3'UTR and trinucleotide repeat microsatellites.//2.2e-06:247:64//U08273
- F-HEMBA1003433//Homo sapiens nibrin (NBS) mRNA, complete cds.//1.4e-149:697:99//AF051334
  - F-HEMBA1003447//Homo sapiens chromosome 4 clone B353C18 map 4q25, complete sequence.//1.7e-77:461: 90//AC004066
  - F-HEMBA1003461//Rhodobacter sphaeroides FIIH (fIIH) gene, partial cds, F1II (fIII) and FIIJ (fIIJ) genes, complete cds.//8.6e-08:752:58//U31090
- F-HEMBA1003463//Homo sapiens chromosome 17, clone HClT305D20, complete sequence.//0.089:172:68// AC004098
  - F-HEMBA1003480//Homo sapiens clone NH0523H20, complete sequence.//4.5e-150:562:97//AC005041
  - F-HEMBA1003528//Streptomyces fradiae gene for trypsinogen precursor, complete cds.//4.7e-09:433:60//D16687 F-HEMBA1003531/Homo sapiens PAC clone DJ1185I07 from 7q11.23-q21, complete sequence.//2.3e-48:297: 90//AC004990
- F-HEMBA1003538//Human complement C1r mRNA, complete cds.//4.3e-22:474:63//M14058
  - F-HEMBA1003545//Rattus norvegicus (clone 1.6kB) islet-2 mRNA, complete cds.//3.5e-143:805:91/L35571
  - F-HEMBA1003555//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 447E6, WORKING
- 20 DRAFT SEQUENCE //3.4e-58:331:83//AL031724 F-HEMBA1003556//Homo sapiens Xp22-175-176 BAC GSHB-484017 (Genome Systems Human BAC Library) complete sequence, 16, 6e-99-703.84//AC005913
  - F-HEMBA1003560/Bovine GTP-binding regulatory protein gamma-6 subunit mRNA, complete cds.//1.3e-99:587: 89/J/J05071
- 25 F-HEMBA1003568//HS\_3149\_A1\_C04\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3149 Col=7 Row=E, genomic survey sequence.//4.1e-05:389.57//AQ166810
  - F-HEMBA1003569//Homo sapiens BAC clone NH0335J18 from 2, complete sequence.//1.6e-102:669:85// AC005539
- F-HEMBA1003571/IDictyostelium discoideum RegA (regA) gene, complete cds.//0.00033:649:58/I/J60170

  F-HEMBA1003579//Plasmodium faiciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P1, WORKING DRAFT SEQUENCE.//0.00034/6256//MAL037444
  - F-HEMBA1003581//Mouse mRNA for talin.//3.3e-41:181:86//X56123
    - F-HEMBA1003591//Homo sapiens chromosome 16, BAC clone RPCI-11\_192K18, complete sequence.//4.4e-70: 273:94//AC006075
- 35 F-HEMBA1003595//Plasmodium falciparum chromosome 2, section 32 of 73 of the complete sequence.//6.0e-17: 768: 58//AF001395
  - F-HEMBA1003597//Plasmodium falciparum chromosome 2, section 35 of 73 of the complete sequence.//4.0e-09: 777:56//AF001398
- F-HEMBA1003598//Homo sapiens PAC clone DJ0537P09 from 7p11.2-p12, complete sequence.//1.3e-146:692: 98//AC005153
- F-HEMBA1003615//HS\_2010\_A2\_A07\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2010 Col=14 Row=A, genomic survey sequence.//1.1e-22:137:97//AQ226592
  - F-HEMBA1003617//Homo sapiens HRIHFB2157 mRNA, partial cds.//2.4e-169:501:97//AB015344
  - F-HEMBA1003621//Mus musculus PIAS3 mRNA, complete cds.//4.7e-37:165:92//AF034080
- 45 F-HEMBA1003622//Plasmodium falciparum 3D7 chromosome 12 PFYAC336 genomic sequence, WORKING DRAFT SEQUENCE, 5 unordered pieces //o.0024:514:58//AC005139
  - F-HEMBA1003630//CIT-HSP-2168N15.TR CIT-HSP Homo sapiens genomic clone 2168N15, genomic survey sequence.//6.5e-15:358:63//B92984
  - F-HEMBA1003637//Homo sapiens clone RG228D17, WORKING DRAFT SEQUENCE, 2 unordered pieces.//5.0e-21:238:76//AC005077
  - F-HEMBA1003640//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 112K5, WORKING DRAFT SEQUENCE.//2.3e-15:371:63//Z85987
    - F-HEMBA1003645//A.thaliana 81kb genomic sequence.//1.0:529:57//X98130
    - F-HEMBA1003646
- 55 F-HEMBA1003656

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- F-HEMBA1003662//Homo sapiens chromosome 17, clone hRPK.332\_H\_18, complete sequence.//1.6e-175:824:
  - F-HEMBA1003667//Homo sapiens chromosome 12p13.3, WORKING DRAFT SEQUENCE, 21 unordered piec-

es.//1.1e-24:190:87//AC004765

F-HEMBA1003679//Homo sapiens BAC clone RG114B19 from 7q31.1, complete sequence.//1.7e-162:579:99//

F-HEMBA1003680//H.sapiens DNA sequence.//7.3e-22:172:87//Z22322

F-HEMBA1003684//H.sapiens mRNA for Miz-1 protein.//0.0054:146:70//Y09723

F-HEMBA1003690//Homo sapiens antigen NY-CO-9 (NY-CO-9) mRNA, partial cds.//2.9e-72:606:77//AF039691 F-HEMBA1003692

F-HEMBA1003711//Homo sapiens chromosome 17, clone HRPC41C23, complete sequence.//0.55:450:60// AC003101

F-HEMBA1003714

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F-HEMBA1003715//Human DNA sequence from clone 931E15 on chromosome Xq25. Contains STSs, GSSs and genomic marker DXS8098, complete sequence.//3.0e-16:316:68//AL023575

F-HEMBA1003720//Homo sapiens chromosome 4 clone B227H22 map 4q25, complete sequence.//1.3e-41:483: 73//AC004056

F-HEMBA1003725//CIT-HSP-2351H9.TF CIT-HSP Homo sapiens genomic clone 2351H9, genomic survey sequence./1.1e-112:532:99//AQ079348

F-HEMBA1003729//HS\_3043\_A1\_E07\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3043 Col=13 Row=I, genomic survey sequence.//1.6e-12:87:98//AQ129345

F-HEMBA1003733//Homo sapiens, clone hRPK.15\_A\_1, complete sequence.//4.7e-104:761:82//AC006213

F-HEMBA1037342/HS, 3027\_A2\_B02\_MF CIT Approved Human Genomic Sperm Library D Homo saplens genomic clone Plate-3027 Col=4 Row=C, genomic survey sequence.//3.4e-0.867.97/I/AQ154731 F-HEMBA1003758/I/T-HSP-2379018.TR CIT-HSP Homo saplens genomic clone 2379018, genomic survey se-

quence.//2.9e-10:310:63//AQ113513

F-HEMBA1003760//Mus musculus hypoxia inducible factor three alpha mRNA, complete cds.//6.4e-114:714:86// AF060194

F-HEMBA1003773//Plasmodium falciparum 3D7 chromosome 12 PFYAC336 genomic sequence, WORKING DRAFT SEQUENCE, 5 unordered pieces.//0.078:378:58//AC005139

F-HEMBA1003783//Human DNA sequence from PAC 509L4 on chromosome 6q22.1-6q22.33. Contains SSX3 like pseudogene, EST, STS.//9.0e-135:804:89//Z99496

30 F-HEMBA1003784//Caenorhabditis elegans cosmid C55B6.//0.054:463:58//U88181

F-HEMBA1003799//Homo sapiens Chromosome 22q11.2 Cosmid Clone 105a In DGCR Region, complete sequence.//1.9e-44:425:76//AC000070

F-HEMBA1003803//Cryctolagus cuniculus troponin T cardiac isoform mRNA, 3' end of cds.//0.95:198:62/L40178
F-HEMBA1003804//Homo sapiens chromosome 17, clone hClT.175\_E\_5, complete sequence.//1.2e-138:275:99//
AC04566

F-HEMBA1003805//Mus musculus quaking type I (QKI) mRNA, complete cds.//6.6e-148:753:95//U44940

F-HEMBA1003807//HS-1068-B1-G06-MR abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 278 Col=11 Row=N, genomic survey sequence.//6.7e-07:241:67//B47212

F-HEMBA1003827//Homo sapiens mRNA for KIAA0616 protein, partial cds.//1.0e-83:586:87//AB014516

40 F-HEMBA1003836//S.cerevisiae chromosome IX cosmid 9150.//5.1e-16:368:63//Z38125

F-HEMBA1003838//CIT-HSP-384J15.TR CIT-HSP Homo sapiens genomic clone 384J15, genomic survey sequence.//1.4e-45:180:90//B54810

F-HEMBA1003856//Homo sapiens chromosome 10 clone CIT9875K-1188B12 map 10p12.1, complete sequence.//0.0014:574:58//AC005875

45 F-HEMBA1003864//, complete sequence.//2.1e-91:234:95//AC005300

F-HEMBA1003866//Mus musculus semaphorin VIa mRNA, complete cds.//5.9e-81:853:71//AF030430

F-HEMBA1003879//H.sapiens CBP80 mRNA.//2.0e-08:87:95//X80030

F-HEMBA1003880//Homo sapiens genomic DNA, chromosome 21q11.1, segment 7/28, WORKING DRAFT SE-QUENCE.//1.7e-180:853:98//AP000036

F-HEMBA1003885//Homo sapiens PAC clone DJ0167F23 from 7p15, complete sequence.//4.5e-39:376:67// AC004079

F-HEMBA1003893//H.sapiens CpG island DNA genomic Mse1 fragment, clone 11b6, forward read cpg11b6.ft1a.// 3.6e-32:173:99//Z59012

F-HEMBA1003902//RPCI11-26M20.TPB RPCI-11 Homo sapiens genomic clone RPCI-11-26M20, genomic survey sequence. I/8.2e-12:422:61//AQ003455

F-HEMBA1003908//Plasmodium falciparum chromosome 2, section 38 of 73 of the complete sequence.//0.0063: 488-58//AF001401

F-HEMBA1003926//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 310O13, WORKING

DRAFT SEQUENCE //3.6e-27:278:76//AL031658

F-HEMBA1003937//Homo sapiens chromosome 3 subtelomeric region.//1.4e-55:315:81//AF109718

F-HEMBA1003939//HS-1047-A1-G04-MF.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 830 Col=7 Row=M, genomic survey sequence.//6.1e-09:413:63//B38195

5 F-HEMBA1003942//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-420 genomic sequence, WORKING DRAFT SEQUENCE. 14 unordered pieces.//0.42:205:65//AC005140

F-HEMBA1003950//M.capricolum DNA for CONTIG MC072.//0.029:458:58//Z33058

F-HEMBA1003953//HS\_2268\_A1\_B04\_MF CIT Approved Human Genomic Sperm Library D Homo saplens genomic clone Plate=2268 Col=7 Row=C, genomic survey sequence.//9.0e-07:239:64//AQ085098

F-HEMBA1003958//Homo sapiens PAC clone DJ0808G16 from 7q11.23-q21, complete sequence.//2.8e-57:424: 74//AC004894

F-HEMBA1003959//RPCI11-78E8.TV RPCI11 Homo sapiens genomic clone R-78E8, genomic survey sequence.//

F-HEMBA1003976//HS\_3146\_A1\_H09\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3146 Col=17 Row=0, genomic survey sequence.//6.3e-10:129:80//AQ141146

F-HEMBA1003978

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F-HEMBA1003985//Caenorhabditis elegans DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone Y105C5, WORKING DRAFT SEQUENCE.//1.0:258:60//Z98855

F-HEMBA1003987

F-HEMBA1003989//Streptomyces coelicolor cosmid 1A9.//0.40:238:61//AL034446

F-HEMBA1004000//Rattus norvegicus satellite seguence d0Mco2.//2.0e-07:116:70//U19354

F-HEMBA1004011//Plasmodium falciparum 3D7 chromosome 12 PFYAC588 genomic sequence, WORKING DRAFT SEQUENCE, 2 unordered pieces.//0.098:286:60//AC004710

F-HEMBA1004012//Homo sapiens chromosome 17, clone hRPK.63\_A\_1, complete sequence.//2.8e-185:896:97//
AC005670

F-HEMBA1004015//Homo sapiens chromosome 17, clone hRPK.721\_K\_1, complete sequence.//6.3e-68:417:80// AC005411

F-HEMBA1004024//Homo sapiens Xp22-83 BAC GSHB-324M7 (Genome Systems Human BAC Library) complete sequence //2 0e-47-418-77//AC005859

30 F-HEMBA1004038//Homo sapiens genomic DNA, chromosome 21q11.1, segment 23/28, WORKING DRAFT SE-QUENCE.//1.6e-51:564:74//AP000052

F-HEMBA1004042//Plasmodium falciparum chromosome 2, section 35 of 73 of the complete sequence.//1.2e-05: 636:55//AE001398

F-HEMBA1004045//Homo sapiens (subclone 1\_g7 from BAC H76) DNA sequence, complete sequence.//1.9e-31: 373:76//AC002252

F-HEMBA1004048//Homo sapiens DNA for P35-related protein, exon 2.//0.039:234:63//D63393

F-HEMBA1004049//Homo sapiens Xp22 GS-524I1 (Genome Systems Human BAC library), complete sequence.//
4.8e-135:780:89//AC003106

F-HEMBA1004055//Human chromosome 3p21.1 gene sequence.//4.7e-09:457:58//L13435

40 F-HEMBA1004056//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 447C4, WORKING DRAFT SEQUENCE.//3.3e-25:246:77//AL021977

F-HEMBA1004074/ICIT-HSP-2053J5.TF CIT-HSP Homo sapiens genomic clone 2053J5, genomic survey sequence./7, 8e-24:233.76/B68555.
F-HEMBA1004086//Saccharomyces doudlasii mitochondrial tRNA-Ser and tRNA-Phe genes, partial sequence.

45 and Var1'p (var1) gene, mitochondrial gene encoding mitochondrial protein, complete cds.//4.5e-08:614:59//
U49822

F-HEMBA1004097//Mus musculus putative transcription factor mRNA, complete cds.//5.9e-121:502:85// AF091234

F-HEMBA1004111//\*\*\* SEQUENCING IN PROGRESS \*\*\* Homo sapiens chromosome 4, BAC clone C0481P14; HTGS phase 1, WORKING DRAFT SEQUENCE, 7 unordered pieces.//2.0e-36:317:80//AC006160

F.-HEMBA1004131/IMMs musculus clone OST2067, genomic survey sequence. IR. 7e-24:320.71/IAF046393
F.-HEMBA1004132/IMS\_3226\_81\_210\_17\_CIT\_Approved Human Genomic Sperm Library D Homo seplens genomic clone Plate=3226 Col=19 Row=H, genomic survey sequence. I/9.7e-13.232:71/IAQ182017

55 F-HEMBA1004138//HS\_3036\_B1\_G11\_MR CIT Approved Human Genomic Sperm Library D Homo saplens genomic clone Plate=3036 Col=21 Row=N, genomic survey sequence.//0.0035:165:64//AQ294763 F-HEMBA1004143

F-HEMBA1004146

- F-HEMBA1004150//Human DNA sequence from PAC 52D1 on chromosome Xq21. Contains CA repeats, STS.// 0.00011:618:60//Z96811
- F-HEMBA1004164//Homo sapiens Xp22-175-176 BAC GSHB-484O17 (Genome Systems Human BAC Library) complete sequence.//2.9e-30:454:68//AC005913
- F-HEMBA1004168//Homo sapiens geminin mRNA, complete cds.//4.5e-133:649:97//AF067855
  - F-HEMBA1004199
  - F-HEMBA1004200//HS\_2015\_A1\_B05\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2015 Col=9 Row=C, genomic survey seguence.//8.5e-34:236:87//AQ247957
- F-HEMBA1004202//Mus musculus chromosome 11, clone mCIT.268\_P\_23, complete sequence.//7.8e-59:216: 83//AC004807
  - F-HEMBA1004203//Homo sapiens clone NH0313P13, WORKING DRAFT SEQUENCE, 15 unordered pieces.//
    6.3e-98:173:98//AC005488
  - F-HEMBA1004207//Homo sapiens leptin receptor short form (db) mRNA, complete cds.//3.2e-166:791:98// U50748
- 15 F-HEMBA1004225//Plasmodium falciparum chromosome 2, section 61 of 73 of the complete sequence.//6.5e-08: 584:60//AE001424
  - F-HEMBA1004227//Rattus norvegicus protein phosphatase 2C mRNA, complete cds.//8.0e-115:713:86//
  - F-HEMBA1004238

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- F-HEMBA1004241/I/CIGSB11 1 check: 4870 from: 1 to: 167234, complete sequence.//0.57:552:58/IAC004708 F-HEMBA1004246//Human DNA sequence from clone 422F24 on chromosome 6q24.1-25.2. Contains a novel gene similar to C. elegans C02C2.5. Contains ESTs, STSs and GSSs, complete sequence.//6.1-e241254:771/
- F-HEMBA1004248//Rattus rattus insulin-induced growth-respons protein (CL-6) mRNA, complete cds.//1.7e-30: 315:74//113610
  - F-HEMBA1004264//Homo sapiens cosmid clone LUCA20 from 3p21.3, complete sequence.//4.4e-07:674:60//
  - F-HEMBA1004267//Homo sapiens chromosome 17, clone hRPC.117\_B\_12, complete sequence.//3.1e-78:335: 87//AC004707
- 30 F-HEMBA1004272//Homo sapiens 12p13.3 PAC RPCI5-1180D12 (Roswell Park Cancer Institute Human PAC Library) complete sequence.//1.4e-176:856:97//AC005831
  - F-HEMBA1004274//HS\_3064\_B2\_A04\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3064 Col=8 Row=B, genomic survey sequence.//3.1e-28:153:100//AQ136993
  - F-HEMBA1004275//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 501A4, WORKING DRAFT SEQUENCE./IS.2e-17:109:99//298051
    F-HEMBA1004276/JCT-185P-2387K6 FT, CIT-HSP Homo saciens genomic clone 2387K6, genomic survey se
    - quence.//5.0e-07:63:98//AQ240477
      F-HEMBA1004286//Homo sapiens TGF beta receptor associated protein-1 mRNA, complete cds.//2.1e-185:868:
    - 99//AF022795
      F-HEMBA1004289//Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone: MQN23, complete sequence.//
  - 1.0:387:59//AB013395
    F-HEMBA1004295/Homo sapiens DNA, anonymous heat-stable fragment RP11-3A.//7.8e-06:92:89//AB012254
    - F-HEMBA1004308/Homo sapiens Idna Bushintos India sequence.//0.00037:413:59//AC004897
      F-HEMBA1004312//Rickettsia prowazekii strain Madrid E, complete genome; segment 2/4.//0.28:522:57//
- 45 AJ235271 F-HEMBA1004321//Homo sapiens chromosome 19, BAC CIT-B-191n6, complete sequence.//7.1e-136:548:92//
  - AC006130
    F-HEMBA1004323//Human DNA sequence from PAC 450C20 on chromosome X.//1.3e-32:320:65//Z84720
  - F-HEMBA1004327//Homo sapiens mRNA for KIAA0522 protein, partial cds.//0.93:222:62//AB011094
- F-HEMBA1004330//Homo sapiens clone DJ1196H06, WORKING DRAFT SEQUENCE, 4 unordered pieces.// 7.0e-168:895:93//AC004995
  - F-HEMBA1004334//Homo sapiens Xp22 BAC 620F15 (Genome Systems BAC library) complete sequence.//4.6e-73;713:75//AC002980
- F-HEMBA1004335//Human DNA-sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 417M14, WORKING
  DRAFT SEQUENCE //1.3e-25:121:85//AL024498
  - F-HFMBA1004341
    - F-HEMBA1004353//\*\*\* ALU WARNING: Human Alu-Sc subfamily consensus sequence.//6.4e-38:278:85//U14571
      F-HEMBA1004354//Human clone C3 CHL1 protein (CHLR1) mRNA, alternatively spliced, complete cds.//4.1e-45:

190:92//U75968

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F-HEMBA1004356

F-HEMBA1004366//P.falciparum complete gene map of plastid-like DNA (IR-A).//2.2e-07:736:57//X95275

F-HEMBA1004372//H. sapiens dystrophin gene intron 44.//1.0:129:62//X77644

F-HEMBA1004389//Mouse interleukin 2 receptor (p55 IL-2R) mRNA, 5' end.//4.7e-42:237:94//M21977

F-HEMBA1004394//Plasmodium falciparum chromosome 2, section 39 of 73 of the complete sequence.//5.2e-05: 519:59//AE001402

F-HEMBA1004396//Human BAC clone RG302F04 from 7q31, complete sequence.//4.0e-32:261:76//AC002463 F-HEMBA1004405//Plasmodium faticiparum 307 tromposome 12 PFYACB8-628 genomic sequence, WORKING DRAFT SE0/URICE 9 unordered pieces //1 4e-07:693-56//AC005507

F-HEMBA1004408//Homo sapiens clone NH0469M07, WORKING DRAFT SEQUENCE, 7 unordered pieces.//
1.2e-69:195:100//AC005037

F-HEMBA1004429//HS\_3193\_A1\_B06\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3193 Col=11 Row=C, genomic survey sequence,//5.1e-67:386:91//AQ172942

F-HEMBA1004433//Human Chromosome 11p11.2 PAC clone pDJ404m15, complete sequence.//3.2e-27:242:82// AC002554

F-HEMBA1004460//Homo sapiens clone DJ0647C14, WORKING DRAFT SEQUENCE, 21 unordered pieces.// 1.7e-75:590:81//AC004846

F-HEMBA1004461//Human DNA sequence from clone 657J8 on chromosome Xq26.1-26.3 Contains GSS, complete sequence.//0.045:215:66//AL034407

F-HEMBA1004479//Mus musculus hypoxia inducible factor three alpha mRNA, complete cds.//5.2e-43:364:79// AF060194

F-HEMBA1004482//Plasmodium falciparum 3D7 chromosome 12 PFYAC181 genomic sequence, WORKING DRAFT SEQUENCE, 8 unordered pieces.//6.8e-17:791:59//AC005505

25 F-HEMBA1004499//Homo sapiens chromosome 17, clone hRPC.1073\_F\_15, complete sequence.//4.4e-125:251: 94//AC004686

F-HEMBA1004502//Plasmodium falciparum 3D7 chromosome 12 PFYAC1122 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered pieces.//0.012:635:57//AC004709

F-HEMBA1004506//Homo sapiens PAC clone DJ0844F09 from 7p12-p13, complete sequence.//2.8e-127:766: 88//4C004453

F-HEMBA1004507

F-HEMBA1004509//Arabidopsis thaliana DNA chromosome 4, BAC clone T10I14 (ESSAII project).//1.0e-13:244: 67//AL021712

F-HEMBA1004534//Human mRNA for actin-binding protein (filamin) (ABP-280).//1.6e-72:678:74//X53416 F-HEMBA1004538//Sequence 1 from patent US 5612190.//0.00015:416:59//i36871

F-HEMBA1004542//Homo sapiens clone NH0486l22, WORKING DRAFT SEQUENCE, 5 unordered pieces.//0.95: 202:64//AC005038

F-HEMBA1004554//Arabidopsis thaliana BAC T26D22.//0.45:624:56//AFO58826

F-HEMBA1004560//Human mRNA for KIAA0281 gene, complete cds.//9.1e-10:173:70//D87457

F-HEMBA1004573//Human BAC clone RG114A06 from 7q31, complete sequence //6.1e-23:134:73i/AC002542 F-HEMBA1004577//Homo sapiens Chromosome 16 BAC clone CIT987SK-582J2, complete sequence //1.6e-15: 190.771/AC004526

F-HEMBA1004586//Homo sapiens clone DJ0810E06, WORKING DRAFT SEQUENCE, 8 unordered pieces.// 3.1e-31:388:76//AC004895

45 F-HEMBA1004596//RPCI11-81021.TJ RPCI11 Homo sapiens genomic clone R-81021, genomic survey sequence.//2.2e-90:458:90//AQ285136

F-HEMBA1004604/Mus musculus COP9 complex subunit 7a (COPS7a) mRNA, complete cds.//8.6e-105:699: 84//AF071316

F-HEMBA1004610//Homo sapiens PAC clone DJ1163J12 from 7q21.2-q31.1, complete sequence.//5.4e-20:267: 72//AC004983

F-HEMBA1004617//CIT-HSP-2319H15.TF CIT-HSP Homo sapiens genomic clone 2319H15, genomic survey sequence.//6.2e-26:147:99//AQ034944

F-HEMBA1004629//Plasmodium falciparum 3D7 chromosome 12 PFYAC1383 genomic sequence, WORKING DRAFT SEQUENCE. 3 unordered pieces.//5.6e-06;766:56//AC005504

F-HEMBA1004631//Human DNA sequence from PAC 368A4 on chromosome X. Contains ESTs, CELLULAR NU-CLEIC ACID BINDING PROTEIN (CNBP) like gene and STSs.//4.7e-73:412:92//Z83843

F-HEMBA1004632//Canine herpesvirus DNA for gene homolog of HSV1 UL16, EHV1 ORF 46, VZV ORF 44.// 0.92:181:61//X90418

- F-HEMBA1004637//G.gallus mRNA for LRP/alpha-2-macroglobulin receptor.//7.8e-47:784:65//X74904
- F-HEMBA1004638//Rattus norvegicus homeodomain protein Nkx6.1 (nkx6.1) mRNA, complete cds.//6.4e-06:458: 61//AF004431
- F-HEMBA1004666//Caenorhabditis elegans DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone Y47D3, WORKING DRAFT SEQUENCE.//0.30:733:55//Z98865
- F-HEMBA1004669//Human DNA sequence from clone 465N24 on chromosome 1p35.1-36.13. Contains two novel genes, ESTs, GSSs and CpG islands, complete sequence.//7.5e-136:521:98//AL031432
  - F-HEMBA1004670//Homo sapiens Chromosome 22q12 Cosmid Clone p90g5, complete sequence.//0.43:365: 59//AC000045
- F-HEMBA1004672

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- F-HEMBA1004693//Plasmodium falciparum 3D7 chromosome 12 PFYAC492 genomic sequence, WORKING DRAFT SEQUENCE, 5 unordered pieces.//0.096:651:54//AC005308
- F-HEMBA1004697//CIT-HSP-2326C13.TR CIT-HSP Homo sapiens genomic clone 2326C13, genomic survey sequence.//0.23:238:65//AQ040642
- 15 F-HEMBA1004705//Homo sapiens Xp22 Cosmid U151G1 (from Lawrence Livermore X library) and PAC RPCI1-93D11 (from Roswell Park Cancer Center) complete sequence.//2.1e-27:375/72//AC002357
  - F-HEMBA1004709//Homo sapiens Xp22-150 BAC GSHB-309P15 (Genome Systems Human BAC Library) complete sequence.//1.6e-36:191:91//AC006210
  - F-HEMBA1004711//Homo sapiens chromosome 17, clone hRPK.271\_K\_11, complete sequence.//1.1e-133:639: 99//AC005562
    - F-HEMBA1004725//RPCI11-75013.TJ RPCI11 Homo sapiens genomic clone R-75013, genomic survey sequence.//6.2e-32:169:100//AQ266512
    - F-HEMBA1004730/Human BAC clone RG035E18 from 7q31, complete sequence.//8.0e-68:732:72//AC004029 F-HEMBA1004733//CIT-HSP-2305M23.TF CIT-HSP Homo sapiens genomic clone 2305M23, genomic survey sequence.//4.0e-18:209-68//AC017556
  - F-HEMBA1004734//Arabidopsis thaliana ubiquitin-conjugating enzyme 17 (UBC17) mRNA, complete cds.//1.8e-13:451:62//AF028340
- F-HEMBA1004736//Human DNA Sequence from PAC 436M11 on chromosome Xp22.11-22.2. Contains the serine threonine protein phosphatase gene PPEF1, and the first coding exon of the RS1 gene for retinoschisis (X-linked, 1997). A Company of the RS1 gene for retinoschisis (X-linked, 1997).
- juvenile) 1 (XLRS1). Contains ESTs, an STS and GSSs, complete sequence.//5.0e-87:646:78//Z94056 F-HEMBA1004748//Human BAC clone RG204I16 from 7g31, complete sequence.//0.24:526:57//AC002461
  - F-HEMBA1004751//Homo sapiens clone DJ0876A24, WORKING DRAFT SEQUENCE, 6 unordered pieces.//
    1.4e-25:268:76//AC004913
  - F-HEMBA1004752//R.norvegicus mRNA for leucocyte common antigen-related protein (3941 bp).//1.1e-07:503: 61//X83546
    - F-HEMBA1004753//Homo sapiens Chromosome 12 Cosmid Clone 6e5, complete sequence.//4.5e-38:314:81//
    - F-HEMBA1004756//Homo sapiens, complete sequence.//1.4e-111:326:84//AC005854
  - F-HEMBA1004758//Sequence 29 from patent US 5534410.//3.9e-135:769:91//I23472
- 40 F-HEMBA1004763//Homo sapiens apoptosis inhibitor survivin gene, complete cds.//3.6e-47:404:79//U75285 F-HEMBA1004768//Homo sapiens PAC clone DJ0979P20 from 7q33-q35, complete sequence.//6.7e-107:890:
  - 78//AC004941
    F-HEMBA1004770//Plasmodium falciparum 3D7 chromosome 12 PFYAC1122 genomic sequence, WORKING
- DRAFT SEQUENCE, 3 unordered pieces.//7.9e-09:806:59//AC004709

  F-HEMBA1004771//G.muris ribosomal RNA operon DNA encoding 16S, 23S and 5.8S ribosomal RNA.//0.69:239:
  - 61//X65063
    - F-HEMBA1004776
    - F-HEMBA1004778
  - F-HEMBA1004795//Drosophila melanogaster A-kinase anchor protein DAKAP550 mRNA, partial cds.//3.4e-46: 778:64//AF003622
  - F-HEMBA1004803//Homo sapiens chromosome Y, clone 264,M,20, complete sequence.//4.3e-82:580:82//
    - F-HEMBA1004806//Homo sapiens BAC clone RG281G05 from 7p15-p21, complete sequence.//5.4e-07:642:59//
- 55 F-HEMBA1004807//Human HIV1 tata element modulatory factor mRNA sequence from chromosome 3.//1.4e-46: 171:92/I/ 01042
  - F-HEMBA1004816//Homo sapiens calpastatin (CAST) gene, exons 10-14.//3.5e-31:546:66//M86257 F-HEMBA1004820//C.botulinum progenitor toxin complex genes.//0.0014:343:62//X87972

- F-HEMBA1004847//Canine mRNA for 68kDA subunit of signal recognition particle (SRP68).//1.5e-85:512:88//
- F-HEMBA1004850//Homo sapiens TGF-beta type I receptor (TGFBR1) gene, exon 1.//0.0065:284:61//AF054590 F-HEMBA1004863//Genomic sequence from Mouse 11, complete sequence.//0.92:250:59//AC000400
- F-HEMBA1004864
  - F-HEMBA1004865//Human DNA sequence from clone 459L4 on chromosome 6p22.3-24.1 Contains EST, STS, GSS, complete sequence.//3.6e-12:214:72//AL031120
  - F-HEMBA1004880//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-319E8, complete sequence.//1.1e-08:255:69//AC004020
- 10 F-HEMBA1004889//Schistocerca americana Antennapedia homeotic protein (Antp) mRNA, complete cds.//0.062: 155:69//U32943
  - F-HEMBA1004900//Plasmodium falciparum unidentified mRNA sequence.//0.00055:323:60//L12043
  - F-HEMBA1004909//Homo sapiens chromosome 17, clone 289A8, complete sequence.//9.6e-16:166:80// AC003051
- F-HEMBA1004918/Turritella communis mitochondrial 16S ribosomal RNA gene, partial./io.81:146.65/M94003 F-HEMBA1004923/Human DNA from overlapping chromosome 19-specific cosmids R32543,, and F15613 containing ZNF gene family member, genomic sequence, complete sequence.//1.4e-58:338.78/IAC030006
  - F-HEMBA1004929//CIT-HSP-2373/16.TR CIT-HSP Homo sapiens genomic clone 2373/16, genomic survey sequence.//2.4e-86:443:96//AQ108676
- F-HEMBA1004930//Homo sapiens PAC clone DJ0608H12 from 7q21, complete sequence.//4.6e-20:219:73// AC004109
- F-HEMBA1004933//HS-1003-A1-E10-MF.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 497 Col=19 Row=I, genomic survey sequence.//1.4e-28:216:85//B30726
- F-HEMBA1004934//Homo sapiens chromosome 21q22.3 PAC 267O10, complete sequence.//0.53:222:61//
  AF042091
  - F-HEMBA1004944//Homo sapiens clone DJ0736H05, WORKING DRAFT SEQUENCE, 5 unordered pieces.//
    1.2e-58:509:78//AC005482
  - F-HEMBA1004954//HS\_2033\_A2\_A08\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2033 Col=16 Row=A, genomic survey sequence.//3.7e-47:243:99//AQ229758
- F-HEMBA1004956//F.falciparum complete gene map of plastid-like DNA (IR-B)./I0.048.421:58//X95276 F-HEMBA1004960//Arabidopsis thailana DNA chromosome 4, ESSA I contig fragment No. 8./I0.89:333:58// 797343
  - E-HEMRA 1004972

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- F-HEMBA1004973//RPCI11-66P8.TK RPCI11 Homo sapiens genomic clone R-66P8, genomic survey sequence.//
- F-HEMBA1004977//Homo sapiens full-length insert cDNA clone YZ83B08.//9.0e-11:84:98//AF086080
  - F-HEMBA1004978/CIT-HSP-2354E10.TR CIT-HSP Homo sapiens genomic clone 2354E10, genomic survey sequence.//0.0021:152:66//AQ075713
    F-HEMBA1004980//HS 3018 A2 E04 MR CIT Approved Human Genomic Sperm Library D Homo sapiens ge-
- nomic clone Plate=3018 Col=8 Row=I, genomic survey sequence.//1.9e-77:392:97//AQ071873
  - F-HEMBA1004983//Albinaria corrugata isolate cor. Prn1.1 16S ribosomal RNA gene, mitochondrial gene for mitochondrial RNA, partial sequence.//0.0030:276:61//AF031680
- F-HEMBA1004995//Homo sapiens chromosome 16, cosmid bridge clone 306E6 (LANL), complete sequence.// 4.2e-138:640.99//AC005590
- 45 F-HEMBA1005008//Human mariner1 transposase gene, complete consensus sequence.//6.8e-20:160:88// U52077
- F-HEMBA1005009//Homo sapiens BAF53a (BAF53a) mRNA, complete cds.//2.0e-144:668:99//AF041474
  F-HEMBA1005019//Homo sapiens mRNA for KIAA0648 protein, partial cds.//1.4e-146:693:98//AB014548
  - F-HEMBA 1000 19/Hollind sapieris Intrivation Knudoven protein; pariaut dus /f. 14e-14h.053/59/Hollind 10496
    F-HEMBA 1000 19/Hollind sapieris DNA sequence from PAC 97D16 on chromosome 6p21.3-22.2. Contains an unknown pseudogene, a 60S Ribosomal protein L24 (1.30) LIKE pseudogene and histone genes H2BFC (H2B/c), H4FFC (H4R) seudogene, 1 H2AFC (H2AC). H3F1K (H3.1 ki) and a RNA-4 in seudogene and RNA-4 Through 104 (1.40) and 1
- Contains ESTs, STSs, GSSs and genomic marker D6S464, complete sequence.//2.2e-115:668:90//AL009179
  F-HEMBA1005035//Homo sapiens chromosome 17, clone hClT.175\_E\_5, complete sequence.//4.6e-138:591:98//
- F-HEMBA1005039//CIT-HSP-2338L5.TR CIT-HSP Homo sapiens genomic clone 2338L5, genomic survey sequence.//3.7e-61:271:88//AQ055486
  - F-HEMBA1005047//Mus musculus mRNA for Rab24 protein.//3.8e-17:218:73//Z22819 F-HEMBA1005050//Human Tis11d gene, complete cds.//0.079:251:63//U07802

- F-HEMBA1005062//Plasmodium falciparum 3D7 chromosome 12 PFYAC69 genomic sequence, WORKING DRAFT SEQUENCE, 4 unordered pieces.//0.018:560:56//AC004688
- F-HEMBA1005066//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 774G10, WORKING DRAFT SEQUENCE.//3 4e-97:432:84//AL034410
- F-HEMBA1005075//H.sapiens DNA 3' flanking simple sequence region clone wg2c3.//6.9e-07:176:68//X76589 F-HEMBA1005079//CIT-HSP-2325M21.TRB CIT-HSP Homo sapiens genomic clone 2325M21, genomic survey sequence (I/2 1e-48:774-93/IA0038720)
  - F-HEMBA1005083//HS\_2248\_B1\_D05\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2248 Col=9 Row=H, genomic survey sequence.//3.4e-06:230:64//AQ129575
- F-HEMBA1005101//Homo sapiens SYT interacting protein SIP mRNA, complete cds.//1.3e-161:762:98// AF080561
  - F-HEMBA1005113//L.esculentum microsatellite repeat DNA region.//0.0038:742:57//X90770
  - F-HEMBA1005123//Homo sapiens clone DJ0673M15, WORKING DRAFT SEQUENCE, 33 unordered pieces.// 9.6e-83:479:78//AC004854
- 15 F-HEMBA1005133//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone Y313F4, WORKING DRAFT SEQUENCE //3.9e-24:576:64//AL023808
  - F-HEMBA1005149//Homo sapiens PAC clone DJ430N08 from 22q12.1-qter, complete sequence.//4.7e-36:283: 80//AC004542
  - F-HEMBA1005152//Homo sapiens chromosome Xp22-67-68, WORKING DRAFT SEQUENCE, 99 unordered pieces.//5.0e-10:332:64//AC004469
  - F-HEMBA1005159//Homo sapiens genomic DNA, chromosome 21q11.1, segment 1/5, WORKING DRAFT SE-QUENCE.//4.0e-10:734:58//AP000023
    - F-HEMBA1005185//H.sapiens CpG island DNA genomic Mse1 fragment, clone 91b2, forward read cpg91b2.ft1a./ 12.2e-14:93:100//Z63847
- 25 F-HEMBA1005201//Drosophila melanogaster cosmid 152A3.//4.7e-35:679:64//AL009194
  - F-HEMBA1005202//Canine mRNA for 68kDA subunit of signal recognition particle (SRP68).//6.7e-138:778:90// x53744
  - F-HEMBA1005206//Drosophila melanogaster Su(P) and anon-73B1 genes and partial o25 gene and Pros26 gene.//7.1e-12:376:62//AJ011320
- F-HEMBA1005219//Homo sapiens mRNA for KIAA0445 protein, complete cds.//7.1e-05:411:60//AB007914
  F-HEMBA1005223//Homo sapiens PAC clone DJ430N08 from 22q12.1-qter, complete sequence.//3.5e-06:212:
  - 66//AC004542
    F-HEMBA1005232//Plasmodium falciparum 3D7 chromosome 12 PFYAC492 genomic sequence. WORKING
- DRAFT SEQUENCE, 5 unordered pieces.//3.7e-07:625:57//AC005308

  5-HEMBA1005241//Homo sapiens PAC clone DJ0777O23 from 7p14-p15, complete sequence.//8.7e-45:567:72//
  AC005154
  - F-HEMBA1005244//Homo sapiens chromosome X clone U177G4, U152H5, U168D5, 174A6, U172D6, and U186B3 from Xp22, complete sequence.//0.96:298:62//AC002365
  - F-HEMBA1005251

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- 40 F-HEMBA1005252//Homo sapiens chromosome 17, clone hRPK.318\_A\_15, complete sequence.//4.5e-160:392: 99//AC005837
  - F-HEMBA1005274//Homo sapiens BAC clone 255A7 from 8q21 containing NBS1 gene, complete sequence.// 2.3e-05:496:60//AF069291
- F-HEMBA1005275//Human DNA sequence from clone 444C7 on chromosome 6p22.3-23. Contains an EST, an STS and GSSs, complete sequence.//5.7e-05:220:64//AL033521
  - F-HEMBA1005293//Homo sapiens echinoderm microtubule-associated protein homolog HuEMAP mRNA, complete cds.//2.4e-20:338:65//U97018
    - F-HEMBA1005296
    - F-HEMB41005304//Human DNA sequence from clone 364l22 on chromosome Xq21.31-22.3. Contains an STS and GSSs, complete sequence//1.6e-51:381:78//AL031012 F-HEMB41005311
      - F-HEMBA1005314//Homo sapiens genomic DNA, chromosome 21q11.1, segment 2/28, WORKING DRAFT SE-QUENCE //0.94-226-63//AP000031
    - F-HEMBA1005315//Homo sapiens BAC810, complete sequence.//9.5e-15:684:62//U85198
- F-HEMBA1005318//Human DNA sequence from PAC 394F12 on chromosome X contains EST, STS, CpG Island clone. I/2.6e-05.472:59//Z83823
  - F-HEMBA1005331//Homo sapiens chromosome 17, clone hRPK.214\_C\_8, complete sequence.//3.3e-90:300:90// AC005803

- F-HEMBA1005338//Homo sapiens mRNA for matrilin-4, partial.//1.4e-151:740:97//AJ007581
- F-HEMBA1005353//CIT-HSP-2310N10.TR CIT-HSP Homo sapiens genomic clone 2310N10, genomic survey sequence.//2.1e-86:438:97//AQ016145
- F-HEMBA1005359//Human zinc finger protein ZNF137 mRNA, complete cds.//1.8e-98:500:88//U09414
- F-HEMBA1005367//Mus musculus melastatin mRNA, complete cds.//8.3e-72:577:73//AF047714
- F-HEMBA1005372//Human DNA sequence from PAC 293E14 contains ESTs, STS.//1.3e-07:274:66//Z82900 F-HEMBA1005374//Homo sapiens clone 277F10, WORKING DRAFT SEQUENCE, 5 unordered pieces.//1.9e-48: 611-88//AC004813
- F-HEMBA1005382//HS\_3063\_B2\_F11\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3063 Col=22 Row=L, genomic survey sequence.//1.6e-27:154:98//AQ103204
  - F-HEMBA1005389//Plasmodium falciparum telomere nucleotide sequence.//4.0e-07;443:61//M23175
    - F-HEMBA1005394//CIT-HSP-2368B11.TR CIT-HSP Homo sapiens genomic clone 2368B11, genomic survey sequence.//7.6e-17:225:71//AQ076749
  - F-HEMBA1005403//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 423B22, WORKING DRAFT SEQUENCE://4.5e-131:278:98//AL034379
  - F-HEMBA1005408/HS\_3007\_B2\_G04\_MR CIT Approved Human Genomic Sperm Library D Homo saplens genomic clone Plate-3007 Col=8 Row=N, genomic survey sequence.//8.0e-06:218:66//AQ294366
    - F-HEMBA1005410//Human DNA sequence from cosmid cU120E2, on chromosome X contains Lowe oculocerebrorenal syndrome (OCRL) ESTs and STS.//1.5e-41:432:76//Z73496
    - F-HEMBA1005411

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- F-HEMBA1005423//Homo sapiens cyclin-dependent kinase inhibitor (CDKN2C) mRNA, complete cds.//1.0e-169: 537:99//AF041248
- F-HEMBA1005426
- F-HEMBA1005443//Homo sapiens chromosome 19, BAC CIT-B-191n6, complete sequence.//7.1e-37:260:76//
  AC006130
- F-HEMBA1005447//CIT-HSP-2173N7.TR CIT-HSP Homo sapiens genomic clone 2173N7, genomic survey sequence.//5.0e-133:631:98//B93234
- F-HEMBA1005468//Human DNA sequence from done 20J23 on chromosome Xq26.2-27.2 Contains ras-related C3 botulinum toxin substrate 1 (P21-RAC1) (ras-like protein TC25) EST, CA repeat, STS, CpG island, complete sequence.//f.5e-118:868:83//AL022576
- F-HEMBA1005469//Homo sapiens chromosome 16, P1 clone 96-4B (LANL), complete sequence.//1.2e-179:838: 99//AC005212
  - F-HEMBA1005472//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 228H13, WORKING DRAFT SEQUENCE //3 4e-20:187:74//AL031985
- F-HEMBA1005474//Homo sapiens genomic DNA, chromosome 21q11.1, segment 12/28, WORKING DRAFT SE-QUENCE.//4.1e-22:445:65//AP000041
  - F-HEMBA1005475//CIT-HSP-2322D14.TR CIT-HSP Homo sapiens genomic clone 2322D14, genomic survey sequence.//6.7e-51:269:97//AQ026941
  - F-HEMBA1005497//HS\_3097\_A2\_G05\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3097 Col=10 Row=M, genomic survey sequence.//1.4e-66:345:96//AQ103810
  - F-HEMBA1005500//Homo sapiens PAC clone DJ1093017 from 7q11.23-q21, complete sequence.//5.4e-178:818:
  - F-HEMBA1005506//Mus musculus (clone 0EBF17) early B-cell factor (EBF) mRNA, complete cds.//2.6e-06:73:
- 45 F-HEMBA1005508//Homo sapiens clone hRPK.1\_A\_1, complete sequence.//0.00012:455:60//AC006196
  - F-HEMBA1005511//Homo sapiens MHC class 1 region.//3.3e-43:421:77//AF055066
  - F-HEMBA1005513//Drosophila melanogaster males-absent on the first (mof) gene, complete cds.//2.3e-20:352: 69//U71219
  - F-HEMBA1005517//Homo sapiens DNA for (CGG)n trinucleotide repeat region, isolate E7.//2.5e-08:431:62// AJ001216
  - F-HEMBA1005518//M.musculus mRNA for paladin gene.//8.2e-90:651:81//X99384
    - F-HEMBA1005520//Homo sapiens clone DJ0876A24, WORKING DRAFT SEQUENCE, 6 unordered pieces.// 7 8e-167:755:99//AC004913
- F-HEMBA1005526//Homo sapiens chromosome 9, clone hRPK.202\_H\_3, complete sequence.//2.4e-42:475:73//
  55 4C006241
- F-HEMBA1005528//Mus musculus mCAF1 protein mRNA, complete cds.//1.2e-94:512:92//U21855
  - F-HEMBA1005530
  - F-HEMBA1005548//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 970A17, WORKING

DRAFT SEQUENCE //9.4e-87:422:99//AL034431

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- F-HEMBA1005552//Homo sapiens PAC clone DJ0807C15 from 7q34-q36, complete sequence.//6.1e-41:486:68// AC004743
- F-HEMBA1005558//Drosophila melanogaster DNA sequence (P1 DS00837 (D87)), complete sequence.//2.9e-19: 306:68//AC004377
  - F-HEMBA1005568//Plasmodium falciparum 3D7 chromosome 12 PFYAC812 genomic sequence, WORKING DRAFT SEQUENCE, 8 unordered pieces.//0.0093;345:60//AC004153
  - F-HEMBA1005570//Plasmodium falciparum chromosome 2, section 44 of 73 of the complete sequence.//4.2e-09: 592:59//AE001407
- F-HEMBA1005576//Homo sapiens mRNA for KIAA0463 protein, partial cds.//5.9e-127:610:98//AB007932 F-HEMBA1005577//HS-1004-A1-E11 -MR abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 498 Col-27 Rowel, cenomic survey sequence.//0.00034:254:64//830971
  - F-HEMBA1005581//Rattus norvegicus mRNA for MEGF5, complete cds.//4.0e-57:826:65//AB011531
- F-HEMBA1005582/HS\_3242\_A1\_B07\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3242 Col=13 Row=C, genomic survey sequence.//1.fe-13:91:98//AQ211275 F-HEMBA1005589
  - F-HEMBA1005588//Homo sapiens PAC clone DJ1188N21 from 7q11.23-q21.1, complete sequence.//8.7e-31:283: 75//AC006025
  - F-HEMBA1005593//Homo sapiens chromosome 17, clone hRPK.332\_H\_18, complete sequence.//8.3e-158:748: 99//AC005746
  - F-HEMBA1005595//CIT-HSP-2309F14.TF CIT-HSP Homo sapiens genomic clone 2309F14, genomic survey sequence //6 4e-30:194:91//A0016527
    - F-HEMBA1005606//CIT-HSP-2326i6.TR CIT-HSP Homo saplens genomic clone 2326i6, genomic survey sequence.//0.0014:132:70//AQ041484
- F-HEMBA1005609/Homo sapiens clone RG315H11, WORKING DRAFT SEQUENCE, 5 unordered pieces.//2.9e-33:249.851/kC005089
  F-HEMBA1005616/Homo sapiens DNA sequence from PAC 43C13 on chromosome Xo21.1-Xo21.3. rab proteins
  - gerany/gerany/transferase component A 1 (rab escort protein 1) (REP-1) (choroideraemia protein) (TCD protein).// 6.5e-29:279-69//AL009175
- 39 F-HEMBA1005621/H-lman DNA sequence "SEQUENCING IN PROGRESS \*\*\* from clone 330012, WORKING DRAFT SEQUENCE./l6.4e-90.158:87/IAL031731 F-HEMBA1005627/IRPC111-34P9 TJ RPC1-11 Homo sapiens genomic clone RPCI-11-34P9, genomic survey se
  - quence.//0.014/168.674.AQ045110

    F-HEMBA1005631/Ir/Ir/Momo sapiens PAC clone DJ1086D14, complete sequence.//1.0e-149:736:93//AC004460
    - F-HEMBA1005632
      F-HEMBA1005634//Human DNA sequence from PAC 187N21 on chromosome 6p21.2-6p21.33. Contains ESTs.//
    - 6.6e-38:452:67//Z98036 F-HFMBA 1005666
- F-HEMBA1005670//Homo sapiens PAC clone DJ0665C04 from 7p14-p13, complete sequence.//5.1e-59:687:74//
  40 AC004850
  - F-HEMBA1005679/Homo sapiens clone DJ0425102, WORKING DRAFT SEQUENCE, 5 unordered pieces.//1.0e-4757:85/IAC005478 F-HEMBA1005680
- F-HEMBA1005685//RPCI11-23D19.TKBR RPCI-11 Homo sapiens genomic clone RPCI-11-23D19, genomic survey sequence.//0.99:228:63//AQ013742
  - F-HEMBA1005699//Human ligand for eph-related receptor tyrosine kinases (EPLG8) mRNA, complete cds.//1.4e-72:406:92//U57001
  - F-HEMBA1005705//Human (D21S172) DNA segment containing (CA) repeat.//0.00040:190:66//X56513
  - F-HEMBA1005717//Plasmodium falciparum MAL3P1, complete sequence.//0.0099:260:63//Z97348 F-HEMBA1005732//Human mRNA for KIAA0003 gene, complete cds.//8.1e-19:151:88//D14697
  - F-HEMBA1005737//Homo sapiens PAC clone DJ1099C19 from 7q21-q22, complete sequence.//5.6e-15:157:79// AC005156
  - F-HEMBA1005746//RPCI11-63N8.TK RPCI11 Homo sapiens genomic clone R-63N8, genomic survey sequence.//
    1.3e-18:113:100//AO238535
- 55 F-HEMBA1005755/Homo sapiens DNA sequence from PAC 95C20 on chromosome Xp11.3-11.4. Contains STSs and the DXS7 locus with GT and GTG repeat polymorphisms, complete sequence.//3 6e-56:764/70//297181 F-HEMBA1005765/Human DNA sequence from PAC 288L1 on chromosome 22q12-qter contains ESTs and polymorphic CA repeat (DZ2S1152)/l/1.1e-30/2757/TI/ZB2196

- F-HEMBA1005780//RPCI11-74E19.TJ RPCI11 Homo sapiens genomic clone R-74E19, genomic survey sequence.//0.0011:283:62//AQ268432
- F-HEMBA1005813//Homo sapiens PAC clone DJ0167F23 from 7p15, complete sequence.//0.14:326:61// AC004079
- F-HEMBA1005815//M.musculus mRNA for skeletal muscle-specific calpain.//6.3e-10:706:59//X92523
  - F-HEMBA1005822//Mouse Bac 291G16, WORKING DRAFT SEQUENCE, 19 unordered pieces.//0.87:417:56// AC003020
    - F-HEMBA1005829//Homo sapiens Chromosome 22q11.2 Fosmid Clone f39e1 In DGCR Region, complete sequence.//8.8e-42:370:79//AC000094
- F-HEMBA1005834//Human DNA sequence from clone 353H6 on chromosome Xq25-26.2. Contains the alternatively spliced SMARCA1 gene for SW1/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 1 (SNF2L1) and a 40S Ribosomal Protein S26 pseudogene. Contains ESTs, STSs and GSSs, complete sequence //2 1e-42/690 67/I/AI 02/2577
  - F-HEMBA1005852//F.rubripes GSS sequence, clone 163A22aE9, genomic survey sequence.//4.3e-07:253:59// AL018749
  - F-HEMBA1005853//CIT-HSP-2289L23.TR CIT-HSP Homo sapiens genomic clone-2289L23, genomic survey se
    - quence.//2.2e-68:333:99//B98952 F-HEMBA1005884//Homo sapiens chromosome 5, BAC clone 78c6 (LBNL H191), complete sequence.//1.9e-57:
- 331:87//AC005351

  F-HEMBA1005891//Homo sapiens PAC clone DJ0997N05 from 7q11.23-q21.1, complete sequence.//5.1e-182:
  - 864:98//AC004945
    - F-HEMBA1005894//Homo sapiens, WORKING DRAFT SEQUENCE, 52 unordered pieces.//3.0e-44:340:80// AC004086
- F-HEMBA1005909/Homo sapiens DNA sequence from PAC 127D3 on chromosome 1q23-25. Contains FMC2 and FMC3 genes for Flavin-containing Monooxygenase 2 and Flavin-containing Monooxygenase 3 (Directly-aniline Monooxygenase (N-Oxide 3, EC1.14.13.8, Dimethylaniline Oxidase 3, FMD II, FMC 3), and a gene for another, unknown, Flavin-containing Monooxygenase family protein. Contains ESTs and GSSs, complete sequence (fl. 3a-12/82/857/IJA) 021026.
  - F-HEMBA1005911//Human DNA sequence from clone 1158E12 on chromosome Xp11.23-11.4 Contains EST, STS, GSS, CpG island, complete sequence //1.0e-44;328;77//AL031584
- STS, GSS, CpG island, complete sequence.//1.0e-44:328:77//AL031584
  F-HEMBA1005921//Homo sapiens chromosome 17, clone hRPK.112 H 10, complete sequence.//1.3e-41:431:
  - 77//AC005666
    F-HEMBA1005931//Homo sapiens chromosome 12p13.3 clone RPCI4-761J14. WORKING DRAFT SEQUENCE.
  - 60 unordered pieces.//1.1e-29:394:70//AC006086
- F-HEMBA1005934//Homo sapiens PAC clone DJ1140G11 from 14q24.3, complete sequence.//8.1e-06:115:80// AC004974
  - F-HEMBA1005962//RPCI11-17O15.TV RPCI-11 Homo sapiens genomic clone RPCI-11-17015, genomic survey sequence.//9.5e-36:315:84//B82821
- F-HEMBA1005963//HS\_3055\_A1\_E08\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3055 Col=15 Row=I, genomic survey sequence.//9.3e-73:372:97//AQ147357
  - F-HEMBA1005990//Homo sapiens I-1 receptor candidate protein mRNA, complete cds.//1.3e-149:697:99//
  - F-HEMBA1005991//Plasmodium falciparum chromosome 2, section 45 of 73 of the complete sequence.//6.3e-07: 423:60//AF001408
- 45 F-HEMBA1005999//Homo sapiens chromosome 4 clone C0026P05 map 4P16, complete sequence.//3.8e-09:360: 64//AC005599
- F-HEMBA1006002

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- F-HEMBA1006002
- F-HEMBA1006005//Homo sapiens MLL (MLL) gene, exons 1-3, and partial cds.//4.5e-83:495:90//AF036405
- F-HEMBA1006031
- F-HEMBA1006035

- F-HEMBA1006036//Human (lambda) DNA for immunogloblin light chain.//2.4e-59:652:74//D87009
  - F-HEMBA1006042//Homo sapiens chromosome 10 clone CIT987SK-1057L21 map 10q25, complete sequence.// 2 1e-43:330:7011AC005386
- F-HEMBA1006067//Plasmodium falciparum 3D7 chromosome 12 PFYAC812 genomic sequence, WORKING
  DRAFT SEQUENCE, 8 unordered pieces.//0.11:433:59//AC004153
  - F-HFMBA1006081
    - F-HEMBA1006090//, complete sequence.//4.5e-139:748:92//AC005500
    - F-HEMBA1006091//Homo sapiens gene encoding telethonin, exons 1 to 2, partial.//0.0091:346:62//AJ011098

- F-HEMBA1006100//Homo sapiens chromosome 10 clone CIT987SK-1143A11 map 10q25, complete sequence.// 2 8e-18:180:78//AC005880
- F-HEMBA1006108//Human DNA sequence from clone 889N15 on chromosome Xq22.1-22.3. Contains part of the
- gene for a novel protein similar to X. laevis Cortical Thymocyte-Marker CTX, the possibly alternatively spliced gene for 26S Proteasome subunit p28 (Ankyrin repeat protein), a novel gene and exons 36 through 45 of the COL446 for Collagen Alpha 6(IV). Contains ESTs, STSs, GSSs and a putative CpG island, complete sequence./i0.26:84: 71/IAI 031177
  - F-HEMBA1006121//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 691N24, WORKING DRAFT SEQUENCE.//5.2e-18:147:87//AL031672
- F-HEMBA1006124//CIT-HSP-2355B17.TF CIT-HSP Homo sapiens genomic clone 2355B17, genomic survey sequence.//0.044:225:61//AO058966
  - F-HEMBA1006130//CIT-HSP-386A20.TF CIT-HSP Homo sapiens genomic clone 386A20, genomic survey sequence.//8.8e-07:173:69//B55085
- F-HEMBA1006138//Homo sapiens DNA sequence from PAC 454M7 on chromosome Xq25-26.3. Contains the OCRL1 gene for Lowo Collocerebrorenal Syndrome protein OCRL-1. Contains ESTs, STSs and GSSs, complete sequence //f. 5e-22.164.75/(A) 022162
  - F-HEMBA1006142//, complete sequence.//7.9e-125:586:99//AC005500
    - F-HEMBA1006155//H.sapiens CpG island DNA genomic Mse1 fragment, clone 119b6, forward read cpg119b6.ft1a.//1.0:85:72//Z64428
- F-HEMBA1006158//Homo sapiens transcription factor forkhead-like 7 (FKHL7) gene, complete cds.//1.1e-185: 852-99//AF048693
  - F-HEMBA1006173//striatum enriched phosphatase=protein-tyrosine-phosphatase [rat, striata, mRNA, 2815 nt].//
  - F-HEMBA1006182//Homo sapiens Chromosome 15q26.1 PAC clone pDJ105i19, complete sequence.//1.4e-22: 194:74//AC005318
  - 104.747A0003010

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- F-HEMBA1006198
- F-HEMBA1006235//Homo sapiens clone 24422 mRNA sequence.//2.6e-175:836:98//AF070557
- F-HEMBA1006248//Pinctada fucata mRNA for insoluble protein, complete cds.//8.2e-05:359:61//D86074
- F-HEMBA1006252//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 531H16, WORKING
  DRAFT SEQUENCE.//0.98:397:58//AL031664
- F-HEMBA1006253
  - F-HEMBA1006259//HS\_2231\_A1\_D10\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2231 Col=19 Row=G, genomic survey sequence.//1.2e-11:233:68//AQ152722
  - F-HEMBA1006268//Homo sapiens Xp22-132-134 BAC GSHB-590J15 (Genome Systems Human BAC library) complete sequence.//5.2e-27:156:85//AC004673
  - F-HEMBA1006272//Human endogenous retrovirus gag mRNA.//8.1e-115:847:80//X72791
    - F-HEMBA1006278//Mus musculus poly(A) polymerase VI mRNA, complete cds.//2.1e-57:665:70//U58134
    - F-HEMBA1006283
- F-HEMBA1006284//Streptomyces fradiae tylactone synthase, starter module and modules 1-7, (tylG) gene, complete cds.//9.6e-06:623:60//U78289
  - F-HEMBA1006291//HS\_2208\_A1\_C03\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2208 Col=5 Row=E, genomic survey sequence.//1.2e-13:105:92//A0091804
    - F-HEMBA1006293//Sequence 8 from patent US 5721351.//5.6e-77:580:75//I89415
- F-HEMBA1006309//Caenorhabditis elegans cosmid F01F1//l1.1e-21.420.63//U13070

  45 F-HEMBA1006310//Ratus norvegicus cytosolic sorting protein PACS-1a (PACS-1) mRNA, complete cds.//6.8e-120.748.85//AF0/76183
  - F-HEMBA1006328//Homo sapiens fragile X mental retardation protein (FMR-1) gene (6 alternative splices), complete cds://1.5e-46:485:73/L29074
  - F-HEMBA1006334//HS-1051-B2-F01-MR.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 773 Col=2 Row=L, genomic survey sequence.//0.0032:61:91//B40563
  - F-HBMBA1006344//HS-1009-A2-B02-MF abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 331 Col=4 Row=C, genomic survey sequence.//3.3e-09:218:66//B31420
    - F-HEMBA1006347//Drosophila melanogaster males-absent on the first (mof) gene, complete cds.//1.6e-31:484: 68//J/71219
- F-HEMBA1006349//HS-1054-A1-G06-MR.abl CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 776 Col=11 Row=M, genomic survey sequence.//5.4e-15:95:100//B41671
  - F-HEMBA1006359//Human ZNF43 mRNA.//1.4e-115:823:81//X59244 F-HEMBA1006364//Mouse mRNA for transforming growth factor-beta2.//2.7e-10:247:71//X57413

- F-HEMBA1006377/I/Mus musculus chromosome 7, done 19K5, complete sequence.//3.0e-57:401:81//AC002327 F-HEMBA1006380//CIT-HSP-2172K18.TF CIT-HSP Homo sapiens genomic clone 2172K18, genomic survey sequence.//3.3e-110:525:59/9182570
- F-HEMBA1006381//HS-1045-B2-F10-MF.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 828 Col=20 Row=L, genomic survey sequence.//4.4e-05:163:70//B37813
- F-HEMBA1006398//Homo sapiens 12q24.2 BAC RPCI11-360E11 (Roswell Park Cancer Institute Human BAC Library) complete sequence.//3.8e-62:370:86//AC004806
  - F-HEMBA1006416//Homo sapiens chromosome 5, P1 clone 1041F10 (LBNL H88), complete sequence.//3.7e-15: 157:78//AC005179
- F-HEMBA106419/Human DNA sequence from clone 71L16 on chromosome Xp11. Contains a probable Zinc Finger protein (pseudo)gene, an unknown putative gene, a pseudogene with high similarity to part of antigen Kl-67, a putative Chondroitin 6-Suliotransferase LiKE gene and a KIAA0267 LIKE putative Na(-):H'(-) exchanger protein gene. Contains a predicted CpG island, ESTs, STSs and GSSs and genomic markers DXS1003 and DXS1055, compilete sequence./II.2e.93752.63/JAQ22165
- 15 F-HEMBA1006421//Homo sapiens chromosome 14q24.3 clone BAC270M14 transforming growth factor-beta 3 (TGF-beta 3) gene, complete cds; and unknown genes.//2.4e-41:438:76//AF107885
  - F-HEMBA106424/H-Human DNA sequence from clone 51J12 on chromosome 6q26-27. Contains the 3 part of the alternatively spliced gene for the human orthologs of mouse GNI-7 and GNI-78 (KH Domain RNA Binding proteins) and zebrafish ZKQ-1 (Quaking protein homolog). Contains ESTs, STSs and GSSs, complete sequence.// 0.27:934-84/AI 031781.
  - F-HEMBA1006426//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 292E10, WORKING DRAFT SEQUENCE.//1.7e-50:310:80//293930
    - F-HEMBA1006438//Liverwort Marchantia polymorpha chloroplast genome DNA.//0.051:440:59//X04465
    - F-HEMBA1006445//Felis catus ras p21 (H-ras) mRNA, partial cds.//1.0:238:59//U62088

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- 25 F-HEMBA1006446//Plasmodium falciparum DNA\*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P6, WORKING DRAFT SEQUENCE, //2, 4e-05;702:58//AL031749
  - F-HEMBA1006461//Homo sapiens chromosome 19, cosmid R30676, complete sequence://8.6e-55:409:83// AC004560
  - F-HEMBA1006467//Homo sapiens chromosome 17, clone hRPK.346\_K\_10, complete sequence.//1.0:293:59//
- F-HEMBA1006471//Plasmodium falciparum 3D7 chromosome 12 PFYAC1122 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered pieces.//1.4e-05:731:59//AC004709
  - F-HEMBA1006474//CIT-HSP-2017H3.TF CIT-HSP Homo sapiens genomic clone 2017H3, genomic survey sequence.//5.2e-60:435:83//B54247
- 35 F-HEMBA1006483//Homo sapiens chromosome 5, BAC clone 8e5 (LBNL H167), complete sequence.//2.9e-48: 28f:84//AC004752
  - F-HEMBA1006485//Homo sapiens BAC clone NH0044G14 from 7q11.23-21.1, complete sequence.//0.96:283: 59//AC006031
- F-HEMBA1008486/Irluman DNA sequence from clone 353H6 on chromosome Xq25-26.2 Contains the alternatively spiled SMARCAI giene for SWIYSNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 1 (SNF2L1) and a 40S Ribosomal Protein S26 pseudogene. Contains ESTs, STSs and GSSs, comulete sequence, IT, 38-14/258/SFI/AUGZ2577
  - F-HEMBA1006489//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 467K16, WORKING DRAFT SEQUENCE.//6.6e-11:595:61//AL031283
- 45 F-HEMBA1006492//Homo sapiens chromosome 17, clone hRPK.269\_G\_24, complete sequence.//6.0e-122:337: 100//AC005828
  - F-HEMBA1006494/I/homo sapiens chromosome 7qtelo BAC E3, complete sequence. I/3.8e-23:459:68/I/AF093117
    F-HEMBA1006497/I/HS\_3023\_B2\_H03\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3023 Co1=8 Row=P, genomic survey sequence. I/2.3e-81:433:95/I/A0033446
  - F-HEMBA1006502//H.sapiens 7SL repeat (clones 2-19b).//1.6e-13:86:87//X62364
    - F-HEMBA1006507/I/homo sapiens mRNA for KIAA0666 protein, partial ofs./I/2.2e-139.470:98I/kB014566
      F-HEMBA100652/I/h/tuma BAC clone RG167605 from 7q21, complete sequence /i/3.e2-724.667.1/kC003991
      F-HEMBA1006530/I/h/tuman DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 1018D12, WORK-INS DRAFT SEQUENCE /I/2.9e-2740.66/KJA031560
- F-HEMBA1006535//Plasmodium falciparum DNA\*\*\* SEQUENCING IN PROGRESS\*\*\* from MAL4P1, WORKING DRAFT SEQUENCE.//0.028:599:60//AL034557
  - F-HEMBA1006540//Homo sapiens multi PDZ domain protein MUPP1 (MUPP1) mRNA, complete cds.//1.4e-171: 654:98//AF093419

- F-HEMBA1006546//Human DNA sequence from cosmid 232L22, between markers DXS366 and DXS87 on chromosome X contains ESTs glycerol kinase pseudogene.//3.8e-104:811:80//Z73986
- F-HEMBA1006559//Mus musculus PRAJA1 (Praia1) mRNA, complete cds://4.8e-99:386:82//U06944
- F-HEMBA1006562//Mus musculus PRAJA1 (Praja1) mRNA, complete cos.//4.8e-99:386:82//006944
  F-HEMBA1006562//Human fructose-1,6-biphosphatase (FBP1) gene, exon 1.//0.012:322:60//U21925
- 5 F-HEMBA1006566//Plasmodium falciparum 3D7 chromosome 12 PFYAC1383 genomic sequence, WORKING DRAFT SEQUENCE. 3 unordered pieces.//0.0026;580:58//AC005504
  - F-HEMBA1006569//Ovis aries beta actin mRNA, complete cds.//6.3e-08:231:70//U39357
  - F-HEMBA1006579//CIT-HSP-2380A22.TR CIT-HSP Homo sapiens genomic clone 2380A22, genomic survey sequence.//0.036:250:62//AQ197107
- F-HEMBA1006583//Mycobacterium tuberculosis H37Rv complete genome; segment 143/162.//1.0:225:63// AI 021841
  - F-HEMBA1006595//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 30A23, WORKING DRAFT SEQUENCE //3.6e-50:689:69I/AL022156
  - F-HEMBA1006597//Homo sapiens Chromosome 7 BAC Clone 239c10, WORKING DRAFT SEQUENCE, 9 unordered pieces.//1.9e-42:253:84//AC004166
- dered pieces.//1.9e-42:253:84//AC004166
  F-HEMBA1006612//RPCI11-88F20.TJ RPCI11 Homo sapiens genomic clone R-88F20, genomic survey se
  - quence.//1.1e-51:266:98//AQ286726
    F-HEMBA1006617//HS\_2193\_B2\_H07\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens ge-
- nomic cione Plate=2193 Col=14 Row=P, genomic survey sequence.//1.1e-594.13.85//AC299685 F-HEMBA1006624/Human DNA sequence from clone 406A7 on chromosome 6q23-24. Contains three pseudogenes similar to Elongation Factor 1-Alpha (EF1-1ALPHA, Stain S1), 60S Acidic Ribosomal Protein P1 and NADH-Ubiquinone Oxidoreductaes 15 K0 as ubunil, and part of the Microfuble Associated Protein E-MAP-115 gene. Con
  - tains ESTs, STSs and GSSs, complete sequence.//1.4e-35:257:89//AL023284
    F-HEMBA1006631//Homo sapiens Chromosome 11g23 PAC clone pDJ356d6, complete sequence.//9.6e-112:
- 25 800:83//AC002036 F-HEMBA1006635//Plasmodium falciparum DNA\*\*\* SEQUENCING IN PROGRESS\*\*\* from MAL1P2, WORKING
  - DRAFT SEQUENCE.//0.15:393:58//AL031745
    F-HEMBA1006639//Petromyzon marinus polyadenylate binding protein (PABP) mRNA, complete cds.//9.6e-15:
- F-HEMBA1006643//Homo sapiens clone DJ0902E20, WORKING DRAFT SEQUENCE, 1 unordered pieces.//
  - 6.88:254:85I/AC006148
    F-HEMBA1006648/Mus musculus integrin binding protein kinase mRNA, complete cds.//1.5e-37:108:88//U94479
    F-HEMBA1006682//Homo sapiens chromosome 5. BAC clone 343g16 (LBNL H180), complete sequence.//1.3e-
  - 154:671:96//AC005601 F-HEMBA1006653

- F-HEMBA1006659//Homo sapiens PAC clone DJ0905J08 from 7p12-p14, complete sequence.//5.2e-110:254:93// AC005189
- F-HEMBA1006665/Homo sapiens Xp22 BAC GSHB-590J6 (Genome Systems Human BAC library) complete sequence.//1.4e-14:177:76//AC004554
- 40 F-HEMBA1006674//Homo sapiens mRNA for nucleolar protein hNop56.//5.5e-15:122:90//Y12065
  - F-HEMBA1006676//Homo sapiens chromosome 19, fosmid 37502, complete sequence.//0.098:218:63//
  - F-HEMBA1006682//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 668J24, WORKING DRAFT SEQUENCE.//1.4e-05:719:57//AL034346
- 45 F-HEMBA1006695//Homo sapiens clone DJ0935K16, complete sequence.//3.1e-22:151:78//AC006011
  - F-HEMBA1006696//CITBI-E1-2522D16.TF CITBI-E1 Homo sapiens genomic clone 2522D16, genomic survey sequence.//5.6e-17:324:66//AQ280738
    - F-HEMBA1006708
  - F-HEMBA1006709
- 50 F-HEMBA1006717//Homo sapiens clone GS308H05, WORKING DRAFT SEQUENCE, 6 unordered pieces.//3.3e-08:136:79//AC005537
  - F-HEMBA1006737//Homo sapiens chromosome 17, clone hRPK.269\_G\_24, complete sequence.//5.8e-162:497: 98//AC005828
- F-HEMBA1006744//Homo sapiens Chromosome 11p14.3 PAC clone pDJ1034g4, complete sequence.//7.4e-48: 320:87//AC004796
  - F-HEMBA1006754/Human DNA sequence from PAC 82.111 and cosmid U134E6 on chromosome Xq22. Contains NIK like and Thyroxin-binding globulin precursor (T4-binding globulin, TBG) genes, ESTs and STSs.//4.1e-129: 844-88/7.23850

- F-HEMBA1006758//Homo sapiens chromosome 5, BAC clone 182a8 (LBNL H161), complete sequence.//2.2e-162.766:99//AC005752
- F-HEMBA1006767//Human Xq28 cosmid U247A3 from LLOXNC01 X chromosome library, complete sequence.// 1.2e-19:326:69//U73465
- F-HEMBA1006778/I/Human DNA sequence from clone 80119 on chromosome 6p21.31-22.2 Contains genes and pseudogenes for olfactory receptor-like proteins, STS, GSS, complete sequence./I/1.4e-103:355:87I/AL022727 F-HEMBA1006780I/CIT-HSP-2359P7.TR CIT-HSP Homo sapiens genomic clone 2359P7, genomic survey se-
  - F-HEMBA1006789//nbxb0037/13r CUGI Rice BAC Library Oryza sativa genomic clone nbxb0037/13r, genomic survey sequence //0.00011:288:63//AO290474
    - F-HEMBA1006795//CIT-HSP-2307E3.TF CIT-HSP Homo sapiens genomic clone 2307E3, genomic survey sequence./5.1e-80.420.96//A0020511
    - F-HEMBA1006796//Human clone 23803 mRNA, partial cds.//4.5e-06;202;68//U79298
    - F-HEMBA1006807//Homo sapiens mRNA for SPOP.//1.2e-66:651:73//AJ000644

quence //0.072:147:68//AQ077208

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- 15 F-HEMBA1006821//Homo sapiens chromosome 17, clone hRPC.62\_O\_9, complete sequence.//6.0e-116:541: 99//AC004797
  - F-HEMBA1006824//Homo sapiens chromosome 19, cosmid R29368, complete sequence.//0.40:159:66// AC004262
  - F-HEMBA1006832//Homo sapiens (subclone 3\_g8 from P1 H25) DNA sequence, complete sequence.//1.8e-24: 323:71//AC002196
    - F-HEMBA1006849//Homo sapiens genomic DNA of 9q32 anti-oncogene of flat epitherium cancer, segment 4/10.// 0.15:403:60//AB020872
    - F-HEMBA1006865//Plasmodium falciparum chromosome 2, section 6 of 73 of the complete sequence.//0.20:472: 57//AF001369
- 25 F-HEMBA1006877//Mus musculus clone OST9241, genomic survey sequence.//3.4e-79:641:76//AF046757
  - F-HEMBA1006885//HS\_2208\_B2\_G06\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2208 Col=12 Row=N, genomic survey sequence.//4.9e-18:206:76//AQ089246
  - F-HEMBA1006900//Human DNA sequence from clone 496N17 on chromosome 6p11.2-12.3 Contains EST, GSS, complete sequence.//5.4e-07:298:65//AL031321
- 30 F-HEMBA1006914//S.pombe chromosome II cosmid c16H5.//0.00040:194:66//AL022104
  - F-HEMBA1006921//Homo sapiens BAC clone GS114I09 from 7p14-p15, complete sequence.//1.1e-174:813:99// AC006027
    - F-HEMBA1006926//Caenorhabditis elegans cosmid ZK185.//0.0075:183:65//AF036704
  - F-HEMBA1006929//P.falciparum complete gene map of plastid-like DNA (IR-A).//4.0e-06:739:57//X95275
- 35 F-HEMBA1006936
  - F-HEMBA1006938//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P4, WORKING DRAFT SEQUENCE.//1.1e-05:733:57//AL031747
  - F-HEMBA1006941//Homo sapiens mRNA for putative thioredoxin-like protein.//1.3e-90:437:98//AJ010841
- F-HEMBA1006949//Human DNA sequence from PAC 363L9 on chromosome X. contains STS and polymorphic CA repeat //0.67:217:62//782205
  - F-HEMBA1006973//Homo sapiens rab3-GAP regulatory domain mRNA, complete cds.//5.6e-143:740:94//
  - F-HEMBA1006976//cDNA encoding alpha 2 to 3 siglytransferase.//2.8e-101:338:89//E06058
- F-HEMBA1006993//Homo sapiens BAC clone BK085E05 from 22q12.1-qter, complete sequence.//7.1e-31:536: 66//4C003071
  - F-HEMBA1006996//Human DNA sequence from clone J428A131, WORKING DRAFT SEQUENCE://9.5e-07:285:
  - F-HEMBA1007002//Genomic sequence for Arabidopsis thaliana BAC F20N2, complete sequence.//0.99:388:58//
  - F-HEMBA1007017//Sequence 3 from Patent WO9416067.//0.96:220:62//A39358
    - F-HEMBA1007018//G.gallus mRNA for dynein light chain-A.//1.3e-124:838:83//X79088
    - F-HEMBA1007045
    - F-HEMBA1007051//Caenorhabditis elegans cosmid Y57G11C, complete seguence.//0.17;343:60//Z99281
- F-HEMBA1007052//Homo sapiens FSHD-associated repeat DNA, proximal region.//4.3e-67:659:74//U85056
- 55 F-HEMBA1007062//Tubulin gene.//1.0:113:67//A18572
  - F-HEMBA1007066//HS\_3116\_A2\_A03\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3116 Col=6 Row=A, genomic survey sequence,//0.80:214:62//AQ140467
  - F-HEMBA1007073//Homo sapiens 12q13 PAC RPCI1-316M24 (Roswell Park Cancer Institute Human PAC library)

- complete sequence //9.3e-54:519:68//AC004242
- F-HEMBA1007078//CIT-HSP-2318N6.TF CIT-HSP Homo sapiens genomic clone 2318N6, genomic survey sequence.//8.7e-80:387:98//AQ044076
- F-HEMBA1007080

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- F-HEMBA1007085//Streptomyces coelicolor cosmid 7A1.//3.5e-06:496:59//AL034447
  - F-HEMBA1007087//Plasmodium falciparum MAL3P6, complete seguence, I/7, 4e-07:553:56//Z98551
  - F-HEMBA1007112//HS\_2171\_A1\_B01\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2171 Col=1 Row=C, genomic survey sequence.//1.0:172:61//AQ091865
- F-HEMBA1007113//Human DNA sequence from clone 1044O17 on chromosome Xp11.3-11.4 Contains GSS and STS, complete sequence //0.54:502:56//AL023875
  - F-HEMBA1007121//Caenorhabditis elegans cosmid ZK430.//1.4e-08:265:64//U42833
    - F-HEMBA1007129//CITBI-E1-2504A5.TF CITBI-E1 Homo sapiens genomic clone 2504A5, genomic survey sequence.//0.97:267:62//AQ264035
  - F-HEMBA1007147//HS\_3208\_A2\_C04\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3208 Col=8 Row=E, genomic survey sequence //9.1e-90:466:95//AQ176696
- F-HEMBA1007149//Homo saplens chromosome 19, cosmid F23149, complete sequence.//6.0e-138:524:98//
  - F-HEMBA1007151//CITBI-E1-2522H6.TF CITBI-E1 Homo sapiens genomic clone 2522H6, genomic survey sequence.//2.0e-20:157:87//AQ280780
- F-HEMBA1007174//Homo sapiens epsin 2a mRNA, complete cds.//2.0e-62:318:97//AF062085
  - F-HEMBA1007178//Homo sapiens chromosome 12p13.3 clone RPCI11-372B4, WORKING DRAFT SEQUENCE, 129 ordered pieces.//1.6e-21:205:80//AC005911
    - F-HEMBA1007194//HS\_3124\_B2\_H08\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3124 Col=16 Row=P, genomic survey sequence,//1.3e-11:87:96//AQ187492
- F-HEMBA1007203//Homo sapiens mRNA for K/AA0214 protein, complete cds.//1.7e-156:478:98//D86987 F-HEMBA1007206//Homo sapiens chromosome 17, clone HRPC837J1, complete sequence.//0.024:342:63//AC004223
  - F-HEMBA1007224//Homo sapiens mRNA for KIAA0797 protein, partial cds.//5.0e-176:839:98//AB018340
  - F-HEMBA1007243//Chinese hamster hprt mRNA, complete cds.//4.3e-58:687:68//J00060
- 30 F-HEMBA1007251//Rabbit troponin T messenger fragment (aa 49 to 129).//0.084:177:62//V00899
  - F-HEMBA1007256//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 328E19, WORKING DRAFT SEQUENCE.//1.3e-75:490:88//AL022240
  - F-HEMBA1007267//HS\_3218\_A1\_F07\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3218 Col=13 Row=K, genomic survey sequence.//2.9e-62:393:87//AQ181128
  - F-HEMBA1007273//CIT-HSP-2171B10.TF CIT-HSP Homo sapiens genomic clone 2171B10, genomic survey sequence.//1.1e-63:314:99//B95401
    - F-HEMBA1007279//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-116A10, complete sequence.//3.1e-31:401:72//AC004638
- F-HEMBA1007281//HS\_3115\_A1\_A11\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3115 Col=21 Row=A, genomic survey sequence.//5.0e-70:372:96//AQ186691
  - F-HEMBA1007288//Human DNA sequence from clone 422G23 on chromosome 6q24 Contains EST, STS, GSS, CpG island, complete sequence.//1.2e-152:727:98//AL031003
    - F-HEMBA1007300//Canis familiaris PDE5 mRNA for 3',5'-Cyclic GMP Phosphodiesterase, complete cds.//2.1e-21:542:63//AB008467
- 45 F-HEMBA1007301//COL1A1=type I collagen pro alpha 1(I) chain propeptide (3' region) [human, fetal cells 86-237, 86-146, 88-251, mRNA Partial Mutant, 855 nt].//1.7e-08:388:61//S64596
  - F-HEMBA1007319//Genomic sequence from Mouse 9, complete sequence.//6.0e-84:390:75//AC000399
  - F-HEMBA1007320
  - F-HEMBA1007322//Homo sapiens BAC clone RG118E13 from 7p15-p21, complete sequence.//0.091:260:64// AC004485
  - F-HEMBA1007327//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-420 genomic sequence, WORKING DRAFT SEQUENCE, 14 unordered pieces.//0.12:472:59//AC005140
  - F-HEMBA1007341//Homo sapiens chromosome 17, clone hRPK.346\_K\_10, complete sequence.//1.5e-18:408: 64//AC006120
- 55 F-HEMBA1007342//Homo sapiens clone DJ1136G02, WORKING DRAFT SEQUENCE, 4 unordered pieces.// 8.7e-25:500:62//AC005377
  - F-HEMBA1007347//Homo sapiens chromosome 5, BAC clone 7g12 (LBNL H126), complete sequence.//0.75:269: 61//AC005738

- F-HEMBB1000005//Homo sapiens chromosome Y, clone 264,M,20, complete sequence.//5.0e-05:441:60// AC004617
- F-HEMBB1000008//Homo sapiens BAC clone RG139P11 from 7q11-q21, complete sequence.//1.0e-44:417:77// AC004491
- F-HEMBB1000018//HS\_2179\_B2\_E04\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2179 Col=8 Row=J, genomic survey sequence.//0.012:87:77//AQ023250
  - F-HEMBB1000024//Human DNA sequence from PAC 106l20 on chromosome 22q12-qter contains NADH pseudogene, ESTs, STS.//8.1e-11:461:61//Z81369
  - F-HEMBB1000025//CIT-HSP-2348F3.TR CIT-HSP Homo sapiens genomic clone 2348F3, genomic survey sequence //0.96:198:62//AQ062938
  - F-HEMBB1000030//Homo sapiens DNA sequence from PAC 32F7 on chromosome X. Contains NUCLEOSOME ASSEMBLY PROTEIN 1-LIKE 3, ESTs.//0.00049:276:64//AL009173
  - F-HEMBB1000036//H.sapiens chromosome 22 CpG island DNA genomic Mse1 fragment, clone 302e2, reverse read 302e2.r.//0.0057:66:81/IZ79857
- F-HEMBB1000037//Homo sapiens erythroblast macrophage protein EMP mRNA, complete cds.//1.9e-100:450: 98//AF084928
  - F-HEMBB1000039//HS\_2167\_B1\_F12\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2167 Col=23 Row=L, genomic survey sequence.//0.022:108:69//AQ092404
- F-HEMBB1000044//borrelia burgdorfari (section 50 of 70) of the complete genome.//1.0e-07/486:61//AE001164
  P-HEMBB1000044//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-628 genomic sequence, WORKING DRAFT SEQUENCE, 9 unordered pieces./f5.3e-05:585:58/IAC005507
  - F-HEMBB1000050//Homo sapiens DNA sequence from clone 501N12 on chromosome 6p22.1-22.3 Contains a gene almost identical to four genes of unknown function, a pseudogene, three (pseudo?) genes similar to genes of unknown function, an unknown gene similar to a rat EST, a PX19 LIKE pseudogene and another unknown gene. Contains ESTs, STSs and GSSs, complete sequence //F.8.6s-38:549/67/JAL022170
  - F-HEMBB1000054//Homo sapiens Xp22 PAC RPCI1-167A22 (from Roswell Park Cancer Center) complete sequence.//7.0e-98:328:83//AC002349
  - F-HEMBB1000055//Homo sapiens genomic DNA for centromeric end of MHC class I region on chromosome 6, cosmid clone: TY2F10, WORKING DRAFT SEQUENCE.//3.7e-05:600:58//AB000880
- F-HEMBB1000059//Homo sapiens clone RG339C12, WORKING DRAFT SEQUENCE, 10 unordered pieces.// 1.3e-48:472:78//AC005096
  - F-HEMBB1000083

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- F-HEMBB1000089//Plasmodiumfalciparum DNA\*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P1, WORKING DRAFT SEQUENCE.//0.0036:679:56//AL031744
- F-HEMBB1000099//Homo saplens chromosome 18 BAC RPCI11-128D14 (Roswell Park Cancer Institute Human BAC Library) complete sequence.//1.1e-15:312:68//AC005909
  - F-HEMBB1000103//Homo sapiens Xp22-150 BAC GSHB-309P15 (Genome Systems Human BAC Library) complete sequence.//1.0e-37:316:74//AC006210
- F-HEMBB1000113//Homo sapiens chromosome 21q22.3 cosmid Q11M15, complete sequence.//3.1e-25:259:76//
  40 AF045450
  - F-HEMBB1000119//Homo sapiens ASMTL gene.//1.2e-137:654:98//Y15521
    - F-HEMBB1000136//Mycobacterium tuberculosis H37Rv complete genome; segment 127/162.//0.59:217:66// Z74697
- F-HEMBB1000141//Homo sapiens DNA from choromosome 19q13.1 cosmid f14121 containing ATP4A and GADPH-2 genes, genomic sequence.//8.4e-31:113:88//AD000090
  - F-HEMBB1000144//Human BAC clone RG114A06 from 7q31, complete sequence.//l.4.e-58:3398.87//AC002542 F-HEMBB1000173//Homo sapiens 12q24 BAC RPCII1-162P23 (Roswell Park Cancer Institute Human BAC library) complete sequence.//9.4e-160:562:93//AC002996 F-HEMBB1000175
  - F-HEMBB1000198//HS\_3071\_A2\_A10\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3071 Col=20 Row=A, genomic survey sequence.//0.99:261:61//AQ137388
    - F-HEMBB1000215//Homo sapiens chromosome 17, clone hRPK.481\_C\_4, complete sequence.//6.7e-17:138:86// AC005839
- F-HEMBB1000217//Arabidopsis thaliana ubiquitin activating enzyme (UBA1) gene, complete cds.//0.00083:287: 60//U80808
- F-HEMBB1000218//Caenorhabditis elegans cosmid C52A11, complete sequence.//0.90:337:56//Z46792
  - F-HEMBB1000226//Human DNA sequence from cosmid RJ14 from a contig from the tip of the short arm of chromosome 16, spanning 2Mb of 16p13.3. Contains ESTs and CpG island.//1.7e-90:175:92//Z69890

- F-HEMBB1000240//Human G-protein-coupled inwardly rectifying potassium channel (KCNJ3) gene, polymorphic repeat sequence.//0.16:171:62//U07918
- F-HEMBB1000244//Homo sapiens clone DJ1129E22, WORKING DRAFT SEQUENCE, 7 unordered pieces.// 4.8e-08:355:63//AC005522
- F-HEMBB1000250//Homo sapiens protein associated with Myc mRNA, complete cds.//6.6e-155;735;98//
  - F-HEMBB1000258//Human adenosine monophosphate deaminase 1 (AMPD1) gene, exons 1-16.//0.58:396:59//
- F-HEMBB1000264//Human clone C3 CHL1 protein (CHLR1) mRNA, alternatively spliced, complete cds.//4.4e-32: 10 100:100//U75968
  - F-HEMBB1000266//Homo sapiens Xp22 BAC GSHB-433024 (Genome Systems Human BAC library) complete sequence.//3.8e-16:176:78//AC004470
  - F-HEMBB1000272//Plasmodium falciparum chromosome 2, section 6 of 73 of the complete sequence.//0.011:379: 58//AF001369
- 15 F-HEMBB1000274//Arabidopsis thaliana DNA chromosome 4, BAC clone T5K18 (ESSAII project).//0.92:272:61//
  - F-HEMBB1000284//Human Xp22 BAC CT-285I15 (from CalTech/Research Genetics), PAC RPCI1-27C22 (from Roswell Park Cancer Center), and Cosmid U35B5 (from Lawrence Livermore), complete sequence, //0.00071:568:
- 20 F-HEMBB1000307//Human DNA sequence from PAC 29K1 on chromosome 6p21.3-22.2. Contains glutathione peroxidase-like; zinc finger, ESTs, mRNA, STS, tRNAs, olfactory receptor pseudogene. I/3.0e-13:439:65//Z98745 F-HEMBB1000312//Homo sapiens clone GS051M12, complete sequence.//0.031:252:65//AC005007
- F-HEMBB1000317//Fugu rubripes GSS sequence, clone 060J22aE10, genomic survey sequence.//0.00033:173; 65//AL026242
- F-HEMBB1000318//HS 3244 B2 H10 T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3244 Col=20 Row=P, genomic survey sequence.//3.9e-85;438:95//AQ252951 F-HEMBB1000335//Homo sapiens chromosome 18, clone hRPK.24 A 23, complete sequence, //0.63:285:61// AC005968
  - F-HEMBB1000336

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- F-HEMBB1000337//Homo sapiens chromosome 4 clone B208G5 map 4g25, complete sequence.//0.0014:309: 64//AC004051
  - F-HEMBB1000338//HS 3108 A2 F07 T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3108 Col=14 Row=K, genomic survey sequence.//3.8e-09:331:63//AQ140356
  - F-HEMBB1000339//Homo sapiens 12c24 PAC RPCI1-46F2 (Roswell Park Cancer Institute Human PAC library) complete sequence //1 2e-52:295:77//AC002351
  - F-HEMBB1000341
    - F-HEMBB1000343//Plasmodium falciparum MAL3P3, complete sequence.//0.00081:397:61//Z98547
    - F-HEMBB1000354//Human DNA sequence from clone 192P9 on chromosome Xp11.23-11.4. Contains a pseudogene similar to rat Plasmolipin, ESTs and GSSs, complete seguence. //9, 1e-34;596;66//AL020989
- 40 F-HEMBB1000369//Genomic sequence from Human 17, complete sequence.//0.012:298:60//AC002090 F-HEMBB1000374//Human Xp22 contig of 3 PACS (R7-39D12, R7-134G1, R7-185L21) from the Roswell Park
  - Cancer Institute, complete sequence, //9.3e-69:294:89//U96409 F-HEMBB1000376//Human DNA sequence from clone 751H9 on chromosome 6g13. Contains part of an unknown
- gene, ESTs, STSs and GSSs, complete sequence.//3.5e-54:352:88//AL034377 45 F-HEMBB1000391//Trichothecium roseum internal transcribed spacer 1, 5,8S ribosomal RNA gene; and internal
- transcribed spacer 2, complete sequence.//0.011:168:67//U51982 F-HEMBB1000399//Homo sapiens Rad17-like protein (RAD17) mRNA, complete cds.//2.6e-163:762:98//
  - F-HEMBB1000402//Homo sapiens Xq28 BAC PAC and cosmid clones containing FMR2 gene exons 1,2, and 3,
  - complete sequence.//7.7e-15:466:63//AC002368
- F-HEMBB1000404//Homo sapiens mRNA for myosin-IXA.//3.5e-65:324:98//AJ001714
  - F-HEMBB1000420//244Kb Contig from Human Chromsome 11p15.5 spanning D11S1 through D11S25, complete sequence //0.013:399:62//AC001228
- F-HEMBB1000434//Homo sapiens PAC clone 278C19 from 12g, complete sequence.//6.1e-83:571:84//AC004263 F-HEMBB1000438//RPCI11-21E14.TP RPCI-11 Homo sapiens genomic clone RPCI-11-21E14, genomic survey sequence.//0.0030:295:63//B83110
  - F-HEMBB1000441//Homo sapiens Chromosome 22q12 Cosmid Clone II47q11, complete sequence, //2.5e-33;372; 72//AC000035

- F-HEMBB1000449/Human DNA sequence from PAC 296K21 on chromosome X contains cytokeratin exon, deltaaminolevulinate synthase (erythroid); F-aminolevulinic acid synthase, (EC 2.3.1.37). 6-phosphofructo-2-kinase/ fructose 2-6-bisphosphatase (EC 2.7.1.105, EC 3.1.3.46). ESTs and STS.//1.3e-5.153472/IZ63821
- F-HEMBB1000455//Saccharomyces cerevisiae mitochondrion origin of replication (ori6) and oli1 gene, complete cds://0.016;522:58//L36899
  - F-HEMBB1000472
  - F-HEMBB1000480

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- F-HEMBB1000487//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 12803, WORKING DRAFT SEQUENCE.//0.00013:314:64//Z98742
- F-HEMBB1000490//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 1185N5, WORKING DRAFT SEQUENCE.//4.1e-110:529:98//AL034423
  - F-HEMBB1000491//Plasmodium falciparum chromosome 2, section 25 of 73 of the complete sequence.//0.10: 187:65//AE001388
  - F-HEMBB1000493//Human DNA sequence from clone 353H6 on chromosome Xq25-26.2. Contains the alternatively spliced SMARCA1 gene for SW1/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 1 (SNF2L1) and a 40S Ribosomal Protein S26 pseudogene. Contains ESTs, STSs and GSSs, complete sequence./i/3.7-e06637:58/IA.022577
    - F-HEMBB1000510//Homo sapiens chromosome 17, clone hRPK.112\_J\_9, complete sequence.//3.1e-96:737:81//
- F-HEMBB1000518//Homo Sapiens Chromosome X clone bWXD171, WORKING DRAFT SEQUENCE, 1 ordered pieces.//0.00014:163:68//AC004676
  - F-HEMBB1000523//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from contig 3-105, complete sequence.//0.41:349:56//AL010212
- F-HEMBB1000530//H.sapiens mRNA for extracellular matrix protein collagen type XIV, C-terminus.//6.6e-37:138: 96/Y11710
  - F-HEMBB1000550//Homo sapiens genomic DNA of 8p21.3-p22 anti-oncogene of hepatocellular colorectal and non-small cell lung cancer, segment 3/11.//3.9e-56:683:71//AB020860
  - F-HEMBB1000554//Homo sapiens \*\*\* SEQUENCING IN PROGRESS \*\*\*, WORKING DRAFT SEQUENCE.//2.2e-51:282:84//AJ011929
- 30 F-HEMBB1000556//Homo sapiens mRNA for KIAA0750 protein, complete cds.//6.1e-32:537:65//AB018293 F-HEMBB1000564
  - F-HEMBB1000573//Homo sapiens clone RG228D17, WORKING DRAFT SEQUENCE, 2 unordered pieces.//8.2e-33:268-73/(AC005077
- F-HEMBB1000575//Human DNA sequence from clone 323M22 on chromosome 22q13.1-13.2. Contains the 5' part of the human ortholog of chicken P52 and mouse H74, and a novel gene coding for a protein similar to KIAA0173 and worm Tubulin Tyrosine Ligase. Contains ESTs, STSs, GSSs, genomic marker D22S418 and putative CpG islands, complete sequence //5.8e-47.734.66//AL022476
  - F-HEMBB1000586//H.sapiens highly polymorphic microsatellite DNA.//0.030:147:67//X79883
  - F-HEMBB1000589//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-279B10, complete sequence.//6.3e-41:278:83//AC002300
  - F-HEMBB1000591//Homo sapiens Xp22 bins 45-47 BAC GSHB-665N22 (Genome Systems Human BAC Library) complete sequence.//1.1e-182:871:98//AC005184
  - F-HEMBB1000592//Hepatitis C virus genomic RNA, 3' nonstranslated region, partial sequence, clone #19.//0.012: 185.64//AF009074
- 45 F-HEMBB1000593//Homo sapiens chromosome 7q22 sequence, complete sequence.//1.2e-131:353:93// AED53356
  - F-HEMBB1000598//Homo sapiens 12p13.3 BAC RPCI3-488H23 (Roswell Park Cancer Institute Human BAC Library) complete sequence.//9.1e-58:600:72//AC006207
    - F-HEMBB1000623//cDNA encoding Coliolus manganese peroxidase.//0.89:284:62//E12284
  - F-HEMBB1000630//Mus musculus clone NSAT47 nonsatellite RNA sequence.//1.9e-15:129:87//U26231
    - F-HEMBB1000631//Sequence 26 from patent US 5708157.//3.2e-27:180:88//80057
      F-HEMBB1000632//Human mRNA for KIAA0351 gene, complete cds.//1.6e-48:811:65//AB002349
    - F-HEMBB1000637//Homo sapiens clone DJ0425I02, WORKING DRAFT SEQUENCE, 5 unordered pieces.//4.1e-58:649:73//AC005478
- 55 F-HEMBB1000638//HS\_3051\_A1\_G01\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3051 Col=1 Row=M, genomic survey sequence.//0.0032/497:56//AQ155234
  - F-HEMBB1000643//Homo sapiens clone RG228D17, WORKING DRAFT SEQUENCE, 2 unordered pieces.//2.4e-50:791:68//AC005077

- F-HEMBB1000649//Homo sapiens Chromosome 16 BAC clone CIT987-SK502C10, complete sequence.//5.2e-64:775:69//AC003009
- F-HEMBB1000652//Homo sapiens chromosome 10 clone CRI-JC2048 map 10q22.1, WORKING DRAFT SE-QUENCE, 4 unordered pieces://2.7e-52:334:89//AC006186
- F-HEMBB1000665//Human DNA sequence from done 452M16 on chromosome Xq21.1-21.33 Contains capping protein alpha subunit isoform 1 pseudogene, STS, GSS, and CA repeat, complete sequence.//0.0062.426.60// AL024493
  - F-HEMBB1000671//Human DNA sequence from PAC 93H18 on chromosome 6 contains ESTs heterochromatin protein HP1Hs-gamma pseudogene, STS and CpG island.//9.6e-95:399:78//Z84488
- F-HEMBB1000673//HS\_3039\_A2\_C08\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3039 Col=16 Row=E, genomic survey sequence.//3.8e-50:293:92//AQ155121
  - F-HEMBB1000684//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 222E13, WORKING DRAFT SEQUENCE.//8.0e-65 :282:83//Z93241
- F-HEMBB1000693//Homo sapiens neuroan1 mRNA, complete cds.//1.6e-118:575:97//AF040723
- 15 F-HEMBB1000705//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-628 genomic sequence, WORKING DRAFT SEQUENCE, 9 unordered pleces.//8.6e-07:251:61//AC005507
  - F-HEMBB1000706//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 153G14, WORKING DRAFT SEQUENCE.//2.9e-20:434:64//AL031118
  - F-HEMBB1000709//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 994L9, WORKING DRAFT SEQUENCE.//0.26:184:65//AL034554
    - F-HEMBE1000725//Rattus norvegicus GTPase Rab8b (Rab8b) mRNA, complete cds.//1.8e-129:692.93/U53475 F-HEMBE1000725//Human Chromosome 16 BAC clone CIT987SK-A-363E6, complete sequence.//2.7e-40:304: 80/J9/1321
- F-HEMBB1000738/Human Xq28 cosmids U126C1, U142F2, U69B6, U145C10, U169A5, U84H1, U24D12, U80A7, U153E6, L35485, and R7-163A6 containing iduronate 2-sulfatase gene and pseudogene, complete sequence.//8.9e-35.582:63/AF011889
  - F-HEMBB1000749//Homo sapiens chromosome 11 clone CIT-HSP-1337H24, WORKING DRAFT SEQUENCE, 9 unordered pieces //6.2e-46:262:89//AC005849
  - F-HEMBB1000763//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 537K23, WORKING DRAFT SEQUENCE //1.6e-99:316:98//AL034405
  - F-HEMBB1000770//Human DNA sequence from clone 80/19 on chromosome 6p21.31-22.2 Contains genes and pseudogenes for offactory receptor-like proteins, STS, GSS, complete sequence.//0.044:325:60//AL022727 F.I-FMBR1000774
    - F-HEMBB1000781//Sequence 3 from patent US 5753446.//1.2e-92:599:86//AR008277
- F-HEMBB1000789/Homo sapiens mRNA for KIAA0677 protein, complete cds.//9.3e-64:672:71/IAB014577 F-HEMBB1000790/Homo sapiens 12q13.1 PAC RPC11-228P16 (Roswell Park Cancer Institute Human PAC Library) complete sequence.//2.4e-41:460:74/I/AC004801
  - F-HEMBB1000794//HS\_3034\_B2\_D12\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3034 Col=24 Row=H, genomic survey sequence.//1.8e-74:378:97//AQ117099
- 40 F-HEMBB1000807//H.sapiens CpG island DNA genomic Mse1 fragment, clone 39d7, reverse read cpg39d7.rt1a.// 8.5e-14:95:97/758412
  - F-HEMBB1000810//H.sapiens chromosome 22 CpG island DNA genomic Mse1 fragment, clone 303a8, complete read.//3.2e-05:138:71//Z79983
- F-HEMBB1000821//HS\_2168\_B1\_A12\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2168 Col=23 Row=B, genomic survey sequence://0.85:208:60//AQ086361
- F-HEMBB1000822//Human BAC clone GS1131H23 from 5p15.2, complete sequence.//3.0e-06:361:60//AC003015 F-HEMBB1000826//Human BAC clone RG180F08 from 7q31, complete sequence.//1.1e-27:360:69//AC002431 F-HEMBB1000827
  - F-HEMBB1000831

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- F-HEMBB1000835//Human DNA sequence from clone 45l4 on chromosome 6q24.1-24.3. Contains two putative unknown genes. ESTs. STSs and GSSs, complete sequence.//0.00098;234;63//AL023581
  - F-HEMBB1000840//Human Chromosome 11 Cosmid cSRL97a6, complete sequence //4.5e-61:3287.9//U/73649
    F-HEMBB1000848//Homo sapiens DNA sequence from PAC 206D15 on chromosome 1q24. Contains a Reduced Folate Carrier protein (RFC) LIKE one. a mitochondrial ATP Synthetase protein 8 (ATP8. MTATP8) LIKE pseu-
- 55 dogene, an unknown gene and the last exon of the JEM1 gene coding for the Basic-Leucine Zipper nuclear factor JEM-1. Contains ESTs, an STS and a BAC end sequence (GSS), complete sequence.//9.7e-144:809:87// A1021068
  - F-HEMBB1000852//Plasmodium falciparum 3D7 chromosome 12 PFYAC293 genomic sequence, WORKING

DRAFT SEQUENCE, 9 unordered pieces //0.12:492:58//AC004157

F-HEMBB1000870//Plasmodium falciparum 3D7 chromosome 12 PFYAC293 genomic sequence, WORKING DRAFT SEQUENCE. 9 unordered pieces://0.0024;212:67//AC004157

F-HEMBB1000876//Homo sapiens ELISC-1 mRNA, partial cds.//1.5e-32:200:94//AF085351

5 F-HEMBB1000883//HS\_3065\_B2\_C04\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3065 Col=8 Row=F, genomic survey sequence.//0.0017:152:66//AQ137687 F-HFMB11000887

F-HEMBB1000888//CIT-HSP-2329A10.TR CIT-HSP Homo sapiens genomic clone 2329A10, genomic survey sequence.//1.5e-31:172:98//AQ044369

F-HEMBB1000890

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F-HEMBB1000893//Plasmodium falciparum MAL3P2, complete sequence.//9.5e-06:768:56//AL034558

F-HEMBB1000908//Homo sapiens clone DJ11119N05, complete sequence.//4.5e-21:199:82//AC004968

F-HEMBB1000910//Plasmodium falciparum DNA\*\*\* SEQUENCING IN PROGRESS\*\*\* from MAL4P1, WORKING DRAFT SEQUENCE.//0.72:366:59//AL034557

F-HEMBB1000913/HS\_3078\_B1\_C02\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic done Plate=3078 Col=3 Row=F, genomic survey sequence.//9.9e-12:221:63/IAQ144507 F-HEMB1000915/Homo sapiens DNA for (CGG)n trinucleotide repeat region, isolate P4/I/12e-49:252:99/H

AJ001215
F-HEMBB1000917//Homo sapiens chromosome 5, P1 clone 254f11 (LBNL H62), complete sequence.//2.3e-42:

316:76//AC006077

F-HEMBB1000927//Human BDR-2 mRNA for hippocalcin, complete cds.//3.6e-30:528:65/D16593

F-HEMBB1000947//CpG0856B CpIOWAgDNA1 Cryptosporidium parvum genomic, genomic survey sequence.//
0.81:262:62//AQ254493

F-HEMBB1000959//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 34606, WORKING DRAFT SEQUENCE //1.2e-43:454:75//Z84487

F-HEMBB1000973//Mus musculus schlafen2 (Slfn2) mRNA, complete cds.//8.3e-42:458:72//AF099973

F-HEMBB1000975//Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone: MBK5, complete sequence.// 0.98:196:63//AB005234

F-HEMBB1000981

30 F-HEMBB1000985//Homo sapiens chromosome 19, cosmid R29388, complete sequence.//2.9e-06:566:57// AC004476

F-HEMBB1000991//Human DNA sequence from PAC 238J17 on chromosome 6q22. Contains EST and STS.// 0.099:391:57//Z98753

F-HEMBB1000996//Human DNA sequence from BAC 999D10 on chromosome 22q13.3. Contains two BAC endsequences (GSSs).//6.2e-33:227:80//Z94802

F-HEMBB1001004

F-HEMBB1001008//Human Chromosome 16 BAC clone CIT987SK-A-951C11, complete sequence.//4.0e-13:164: 79//AC002551

F-HEMBB1001011//Human Chromosome 16 BAC clone CIT987SK-A-635H12, complete sequence.//7.5e-13:229: 69//AC002310

F-HEMBB1001014//Homo sapiens chromosome 16, BAC clone 375G12 (LANL), complete sequence.//0.32:474:

56/ACCUS/31
F-HEMBB1001020//Homo saplens BAC clone 255A7 from 8q21 containing NBS1 gene, complete sequence.//
2.6e-39:218:80//AF069291

F-HEMBB1001024//Homo sapiens BAC clone 393122 from 8q21, complete sequence J/5.3e-05:656:59//AF070717 F-HEMBB1001037//CIT-HSP-2568K16, TF CIT-HSP Homo sapiens genomic clone 2358K16, genomic survey sequence. J/6.6e-05-226464/AQ080539

F-HEMBB1001047//Homo sapiens cosmids Qc14E2, Qc12H12, Qc11F9, Qc10G9, LA1733 and Qc17B8 from Xc28, complete sequence.//4 (e--27:385:71//U82671

F-HEMBB1001051//H.sapiens mRNA for FAN protein.//1.2e-27:160:98//X96586

F-HEMBB1001056//Homo sapiens clone DJ0953A04, WORKING DRAFT SEQUENCE, 5 unordered pieces.// 2.3e-89:180:91//AC006014

F-HEMBB1001058//Homo sapiens 3p22-8 PAC RPCI4-736H12 (Roswell Park Cancer Institute Human PAC Library) complete sequence.//1.2e-41:468:74//AC006060

55 F-HBMBB1001060//Human Tigger1 transposable element, complete consensus sequence.//4.3e-122:785:86//

F-HEMBB1001063//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 523G1, WORKING DRAFT SEQUENCE./I7.1e-162:770:99//AL034375

- F-HEMBB1001068//Homo sapiens liprin-beta2 mRNA, partial cds.//3.1e-146:736:95//AF034803
- F-HEMBB1001098//Bluchnera aphidicola genomic fragment containing (chaperone Hsp60) groEL, DNA biosynthesis initiating protein (dnaA), ATP operon (alpCDGAHFEB), and putative chromosome replication protein (gldA) genes, complete cds; and termination factor Rho (rho) gene, partial cds //0.00088.690.57/IAF6008210
- F-HEMBB1001102//Homo sapiens huntingtin interacting protein HYPH mRNA, partial cds.//2.1e-76:368:99// ΔΕ049612
  - F-HEMBB1001105//CIT-HSP-2185N1.TR CIT-HSP Homo sapiens genomic clone 2185N1, genomic survey sequence //1 0a-09:136:76//AQ002987
  - F-HEMBB1001112//Rattus rattus sec61 homologue mRNA, complete cds.//1.0e-108:909:76//M96630
- F-HEMBB1001114//Homo sapiens chromosome 17, clone hRPK.795\_F\_17, complete sequence.//7.2e-07:459: 59//AC005284
  - F-HEMBB1001117//HS\_2178\_B1\_E12\_MR CIT Approved-Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2178 Col=23 Row=J, genomic survey sequence./7.8e-50:331:86//AQ068244
  - F-HEMBB1001119//Human collagen type XII alpha-1 precursor (COL12A1) mRNA, complete cds.//1.6e-25:150: 98//U73778
- F-HEMBB1001126

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- F-HEMBB1001133//Homo sapiens Xp22-132-134 BAC GSHB-590J15 (Genome Systems Human BAC library) complete sequence.//2.8e-24:228:80//AC004673
- F-HEMBB1001137
  F-HEMBB1001142//Homo sapiens chromosome Y, clone 264,M,20, complete sequence.//1.0e-40:231:76//
  - AC004617
    F-HEMBB1001151//Rattus norvegicus golgi peripheral membrane protein p65 (GRASP65) mRNA, complete cds.//
  - 2.9e-47:640:67//AF015264
    F-HEMBB1001153/CIT-HSP-2359K11.TR CIT-HSP Homo sapiens genomic clone 2359K11. genomic survey se-
- 25 quence.//0.76:136:67//AQ075724
  - F-HEMBB1001169//Human DNA sequence from PAC 84F12 on chromosome Xq25-Xq26.3. Contains glypican-3 precursor (intestinal protein OCI-5) (GTR2-2), ESTs and CA repeat.//9.9e-63:259:79//AL008712
  - F-HEMBB1001175//Human mRNA for ankyrin motif, complete cds.//2.2e-34:509:66//D78334
    F-HEMBB1001177//CIT-HSP-2321117.TR CIT-HSP Homo sapiens genomic clone 2321117, genomic survey se-
- 30 quence.//5.9e-27:320:75//AQ036473 F-HEMBB1001182//RPCI11-30J5.TV RPCI-11 Homo sapiens genomic clone RPCI-11-30J5, genomic survey se
  - quence.//5.7e-06:62:96/f885188
    F.HFMRR1001199
    - F-HEMBB1001208//HS\_2026\_B1\_C07\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2026 Col=13 Row=F, genomic survey sequence.//0.00018:134:70//AQ229237
- F-HEMBB1001209//CITBI-E1-2521F23.TF CITBI-E1 Homo sapiens genomic clone 2521F23, genomic survey sequence.//1.4e-95.464:98//AQ278357
  - F-HEMBB1001210//HS\_3102\_A2\_F09\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3102 Col=18 Row=K, genomic survey sequence.//2.6e-90:446:98//AQ119196
- 40 F-HEMBB1001218//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from done 796F18, WORKING DRAFT SEQUENCE.//1.0e-31:315.72//AL031291
  - F-HEMBB1001221//Plasmodium falciparum 3D7 chromosome 12 PFYAC1383 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered places.//9.7e-17:770:59//AC005504
- F-HEMBB1001234/H.sapiens CpG island DNA genomic Mse1 fragment, clone 39f9, forward read cpg39f9.ft1e// 4.0e-30:171:97//Z65435
  - F-HEMBB1001242//Homo sapiens mRNA for LAK-1, complete cds.//3.8e-30:458:67//AB005754
    - F-HEMBB1001249//CIT-HSP-2375N19.TF CIT-HSP Homo sapiens genomic clone 2375N19, genomic survey sequence.//0.0076:250:63//AQ109087
  - F-HEMBB1001253//Homo sapiens genomic DNA, chromosome 21q11.1, segment 3/28, WORKING DRAFT SE-QUENCE.//0.0097:89:80//AP000032
  - F-HEMBB1001254//CIT-HSP-2320E5.TF CIT-HSP Homo sapiens genomic clone 2320E5, genomic survey sequence.//3.7e-54:284:97//AQ037173
    - F-HEMBB1001267//Homo sapiens chromosome 17, clone hRPK.488\_L\_1, complete sequence.//3.5e-30:236:78// AC005303
- F-HEMBB1001271//HS\_3011\_A1\_G02\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3011 Col=3 Row=M, genomic survey sequence.//5.2e-07:364:62//AQ214217
  - F-HEMBB1001282//CIT-HSP-2356J20.TF CIT-HSP Homo sapiens genomic clone 2356J20; genomic survey sequence.//1.8e-16:109:97//AQ060969

- F-HEMBB1001288//R.norvegicus mRNA for gephyrin.//3.4e-18:194:77//X66366
- F-HEMBB1001289//Genomic sequence from Human 9q34, complete sequence.//4.8e-66:434:74//AC000387

F-HEMBB1001294//HS\_3039\_B1\_D01\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3039 Col=1 Row=H, genomic survey sequence.//2.0e-90:437:99//AQ155035

F-HEMBB1001302

- F-HEMBB1001304//CIT-HSP-2053E15.TF CIT-HSP Homo sapiens genomic clone 2053E15, genomic survey sequence //2.2e-07:370:61/B69144
- F-HEMBB1001314//Mus musculus Olf-1/EBF-like-3 transcription factor (O/E-3) mRNA, complete cds.//5.7e-116: 663:85//U92703
- F-HEMBB1001315//Homo sapiens chromosome 10 clone LA10NC01\_40\_G\_3 map 10q26.1-10q26.2, WORKING DRAFT SEQUENCE, 1 ordered pieces.//2.5e-33:328:77//AC006096
  - F-HEMBB1001317//Homo sapiens Xp22-150 BAC GSHB-309P15 (Genome Systems Human BAC Library) complete sequence.//1.4e-122:680:91//AC006210
  - F-HEMBB1001326//Homo sapiens BAC clone RG136N17 from 7p15-p21, complete sequence.//2.8e-09:518:60// AC004129
- AC004129 F-HEMBB1001331//Mus musculus mRNA for hepatoma-derived growth factor, complete cds, strain: BALB/c.//3.7efor the page 2016.
  - 5-HEMBB1001335//HS\_3055\_A1\_H10\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3055 Col=19 Row=0, genomic survey sequence.//1.0:222:63//AQ147384
- 20 F-HEMBB1001337/Irluman PAC clone DJ0093/03 from Xq23, complete sequence./I/1.0e-74:319.88/I/AC0039935 F-HEMBB1001339/I/Homo saplens FSHD-associated repeat DNA, proximal region./I/4.0e-135:856:87/I/U85056 F-HEMBB1001346/I/human familial Alzheimer's disease (STM2) gene, complete cds./I/3.3e-44:48174/I/L50871
  - F-HEMBB1001348/Homo sapiens BAC done NH0491B03 from 7p21-p15, complete sequence.//1.8e-17:210:73//
    AC006041
- F-HEMBB1001356//Homo sapiens clone RG252P22, WORKING DRAFT SEQUENCE, 3 unordered pieces.//1.0: 386:59//AC005079
  - F-HEMBB1001364//Homo sapiens chromosome 17, clone hRPC.842\_A\_23, complete sequence.//0.97:349:61// AC004662
  - F-HEMBB1001366//Homo sapiens chromosome 10 clone CIT987SK-1188I5 map 10p11.2-10p12.1, complete sequence.//5.5e-161:766:98//AC005876
    - F-HEMBB1001367//Homo sapiens chromosome 17, clone hRPC.906\_A\_24, complete sequence.//3.0e-55:510: 76//AC004408
    - F-HEMBB1001369//Homo sapiens BAC clone RG163K11 from 7q31, complete sequence.//0.048:244:64// AC005192
- 35 F-HEMBB1001380//Homo sapiens PAC clone DJ1102B04 from 7q11.23-7q21, complete sequence.//2.5e-26:257: 78//AC006204
  - F-HEMBB1001384//Mus musculus COP9 complex subunit 4 (COPS4) mRNA, complete cds.//5.1e-99:571:89// AF071314
- F-HEMBB1001387//Leishmania tarentolae mitochondrial 12S ribosomal RNA gene.//7.1e-05:546:58//X02354

  F-HEMBB1001394//Homo sapiens BAC clone GS421103 from Xq25-q26, complete sequence.//4.0e-129:788:88//
  - F-HEMBB1001410//Homo sapiens wbscr1 (WBSCR1) and replication factor C subunit 2 (RFC2) genes, complete cds.//4.8e-11:632:59//AF045555
- F-HEMBB1001424//Mus musculus Chromosome 4 BAC clone BacB6, complete sequence.//0.0012:435:59//
  45 AC003019
  - F-HEMBB1001426//Homo sapiens clone DJ0736H05, WORKING DRAFT SEQUENCE, 5 unordered pieces.//
    - F-HEMBB1001429//leucine aminopeptidase [cattle, kidney, mRNA, 2056 nt].//4.1e-114:668:88//S65367
    - F-HEMBB1001436//Homo sapiens FUT2 gene, intron 1, complete sequence.//2.3e-37:438:74//AB000931
  - F-HEMBB1001443//Bos taurus pyruvate dehydrogenase phosphatase mRNA, complete cds.//9.1e-92:550:88// L18966
    - F-HEMBB1001449//Homo sapiens chromosome 5, PAC clone 228g9 (LBNL H142), complete sequence.//0.00024: 385:62//AC004768
- F-HEMBB1001454//Homo sapiens chromosome 19, cosmid R34169, complete sequence.//0.84:577:57// AC005790
  - F-HEMBB1001458//Human Chromosome 11 pac pDJ197h17, WORKING DRAFT SEQUENCE, 11 unordered pieces.//8.0e-40:377:78//AC000382
    - F-HEMBB1001463//Human Chromosome 16 BAC clone CIT987SK-A-270G1, complete sequence.//0.011:482:

59//AF001549

F-HEMBB1001464//Human chromosome 16p13 BAC clone CIT987SK-3H8 complete sequence.//0.019:263:61// 1191320

F-HEMBB1001482//Rattus norvegicus Olf-1/EBF associated Zn finger protein Roaz mRNA, alternatively spliced form, complete cds.//1.0e-30:521:66//U92564

F-HEMBB1001500//Homo sapiens clone DJ0742P04, WORKING DRAFT SEQUENCE, 6 unordered pieces.// 1.3e-31:479:71//AC004873

F-HEMBB1001521//Homo sapiens clone RG269P13, WORKING DRAFT SEQUENCE, 6 unordered pieces, //3.7e-51:680:70//AC005080

F-HEMBB1001527

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F-HEMBB1001531//Homo sapiens Chromosome 22q11.2 Cosmid Clone 89h In DGCR Region, complete sequence.//1.3e-79:696:79//AC000089

F-HEMBB1001535//0.aries DNA for polymorphic marker 'OVINRA01' (339 bp).//0.00034:217:62//X89268

F-HEMBB1001536//Homo sapiens PAC clone DJ1182N03 from 7g11,23-g21.1, complete sequence.//0.54:266; 60//AC004548

F-HEMBB1001537//Homo sapiens chromosome 19, cosmid R29368, complete sequence.//4.6e-25:784:61// AC004262

F-HEMBB1001555//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-248F7, complete sequence.//6.9e-50:213:80//AC004605

F-HEMBB1001562//Homo sapiens clone NH0523H20, complete sequence.//0.46:269:60//AC005041

F-HEMBB1001564//Human DNA sequence from clone 192P9 on chromosome Xp11,23-11.4. Contains a pseudogene similar to rat Plasmolipin, ESTs and GSSs, complete sequence.//1.7e-107:620:83//AL020989

F-HEMBB1001565//Homo sapiens BAC clone RG437L15 from 8g21, complete sequence.//2.4e-50:734:67// AC004003

25 F-HEMBB1001585//Human DNA sequence from clone 790B6 on chromosome 20p11.22-12.2. Contains STSs and GSSs, complete sequence.//1.4e-166:816:97//AL031677

F-HEMBR1001586

F-HEMBB1001588//Homo sapiens chromosome 19, CIT-HSP-444n24, complete sequence.//1.6e-21:419:65// AC005261

30 F-HEMBB1001603

> F-HEMBB1001618//Homo sapiens DNA sequence from PAC 142L7 on chromosome 6q21. Contains a Laminin Alpha 4 (LAMA4) LIKE gene coding for two alternatively spliced transcripts, a Tubulin Beta LIKE pseudogene, a Connective tissue growth factor (NOV, GIG) LIKE gene, A predicted CpG island, ESTs, STSs and genomic marker D6S416, complete sequence.//4.5e-29:422:72//Z99289

35 F-HEMBB1001619//HS 3079 B1 A04 MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3079 Col=7 Row=B, genomic survey sequence.//0.0010:77:79//AQ123388 F-HEMBB1001630//Homo sapiens clone RG315H11, WORKING DRAFT SEQUENCE, 5 unordered pieces.//3.2e-

12:667:59//AC005089

F-HEMBB1001635//Plasmodium falciparum MAL3P7, complete sequence.//3.8e-05:475:57//AL034559

40 F-HEMBB1001637//Homo sapiens DNA sequence from PAC 934G17 on chromosome 1p36.21. Contains the alternatively spliced CLCN6 gene for chloride chanel proteins CLC-6A (KIAA0046) -B, -C and -D, the alternatively spliced NPPA gene coding for Atrial Natriuretic Factor ANF precursor (Atrial Natriuretic peptide ANP, Prepronatriodilatin), the NPPB gene for Brain Natriuretic Protein BNP, and a pseudogene similar to SBF1 (and other Myotubularin-related protein genes). Contains ESTs, STSs and the genomic marker D1S2740, complete sequence.// 45 9.2e-13:168:76//AL021155

F-HEMBB1001641//Arabidopsis thaliana genomic DNA, chromosome 5, P1 clone: MPO12, complete sequence.// 0.00097:721:58//AB006702

F-HEMBB1001653//Homo sapiens chromosome 2 clone 101B6 map 2p11, complete sequence.//0.15:276:63// AC002038

F-HEMBB1001665//Bovine herpesvirus type 1 early-intermediate transcription control protein (BICP4) gene, complete cds.//0.43:393:61//L14320

F-HEMBB1001668//F16C15-T7 IGF Arabidopsis thaliana genomic clone F16C15, genomic survey sequence.// 0.040:275:60//B12308

F-HEMBB1001673//Homo sapiens mRNA for KIAA0646 protein, complete cds.//7.2e-171:803:98//AB014546

F-HEMBB1001684//Sequence 1 from patent US 5700927.//7.5e-124:883:81//86429

F-HEMBB1001685//CIT-HSP-2287O9.TF CIT-HSP Homo sapiens genomic clone 2287O9, genomic survey sequence.//2.3e-34:191:97//B99261

F-HEMBB1001695//Human DNA sequence from clone 431P23 on chromosome 6q27. Contains the first coding

exon of the MLLT4 gene for myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 4 (AF-6, Afadin, MLLT-4, ALL-1 fusion partner), and a Serine Palmitoy(transferase 2 (EC 2.3.1.50, Long Chain Base Biosynthesis protein 2, LCB-2, SPT-2) pseudogene. Contains ESTs, STss, GSSs, and a putative CpG island. complete sequence. (IV) 0.091:334-63/JAL.009178

F-HEMBB1001704//Human DNA sequence from clone 931E15 on chromosome Xq25. Contains STSs, GSSs and genomic marker DXS8098, complete sequence.//1.2e-17:144:87//AL023575 F-HEMBB1001706

F-HEMBB1001707//Guinea pig CD19 mRNA, complete cds.//0.57:232:62//M62543

F-HEMBB1001717//Saccharomyces cerevisiae mitochondrial tRNA-Tyr, tRNA-Asn, & amp; tRNA-Met genes.//
1.1e-13:723:58//AJ223323

F-HEMBB1001735//Human PAC clone DJ0596009 from 7p15, complete sequence.//1.3e-36:427:73//AC003074 F-HEMBB1001736//S.pombe chromosome II cosmid c4B4.//0.0085:479:57//AL023706

F-HEMBB1001747//Homo sapiens PAC clone DJ1002N02 from 7p21-p22, complete sequence.//4.0e-112:532: 84//AC005376

F-HEMBB1001749//Homo sapiens chromosome 17, clone hRPK.259\_G\_18, complete sequence.//1.3e-98:395: 82//AC005829

F-HEMBB1001753//S.maximus repeat region, 342bp.//4.2e-11:69:85//Z78099

F-HEMBB1001756//Homo sapiens full-length insert cDNA clone ZD86A11.//0.0015:302:62//AF088064

F-HEMBB1001760//P.falciparum complete gene map of plastid-like DNA (IR-A).//0.011:615:56//X95275

F-HEMBB1001762//CIT-HSP-2290J16.TF CIT-HSP Homo sapiens genomic clone 2290J16, genomic survey sequence.//0.84:208:64//AQ005184

F-HEMBB1001785//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P3, WORKING DRAFT SEQUENCE.//0.0019:469:60//AL031746

F-HEMBB1001797//Human heterogenous nuclear RNA W16W.//0.00012:83:86//X17272

F-HEMBB1001802//Plasmodium falciparum MAL3P7, complete sequence.//1.8e-11:538:60//AL034559

F-HEMBB1001812//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 356B8, WORKING DRAFT SEQUENCE.//1.0e-56:304:84//Z98882

F-HEMBB1001816//Homo sapiens chromosome 19, cosmid F24083, complete sequence.//3.6e-75:300:87// AC005204

30 F-HEMBB1001831//Homo sapiens PAM COOH-terminal interactor protein 1 (PCIP1) mRNA, complete cds.//2.3e-162:763:98//AF056209

 $F-HEMBB1001834//CIT-HSP-2291012.TF\ CIT-HSP\ Homo\ sapiens\ genomic\ clone\ 2291012,\ genomic\ survey\ sequence. \ If T.6e-08:73:94//AQ004168$ 

F-HEMBB1001836//Homo sapiens 12q13.1 PAC RPCI1-228P16 (Roswell Park Cancer Institute Human PAC Library) complete sequence.//5.7e-30:297:79//AC004801

F-HEMBB1001839//Human Chromosome X, complete sequence.//0.016;293;63//AC004073

F-HEMBB1001850//Plasmodium falciparum 3D7 chromosome 12 PFYAC1383 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered pieces.//0.0027:812:58//AC005504

F-HEMBB1001863//Human Chromosome 15q26.1 PAC clone pDJ460g16, WORKING DRAFT SEQUENCE, 3 unordered pieces //8.3e-43:520:72//AC004581

F-HEMBB1001867//Human proto-oncogene tyrosine-protein kinase (ABL) gene, exon 1a and exons 2-10, complete cds.//1.7e-56:399:86//U07563

F-HEMBB1001868//Rattus norvegicus clone 923 polymeric immunoglobulin receptor mRNA 3' untranslated region, GA rich region, and microsatellites with GGA-triplet and GAA-triplet repeats.//6.1e-08:234:67//U01145

45 F-HEMBB1001869//Homo sapiens full-length insert cDNA clone YT86F01.//7.4e-87:432:97//AF085974 F-HEMBB1001872

F-HEMBB1001874//Homo sapiens clone DJ241P17, WORKING DRAFT SEQUENCE, 7 unordered pieces.//3.4e-14:631:61//AC005000

F-HEMBB1001875//Human DNA sequence from clone J428A131, WORKING DRAFT SEQUENCE.//0.93:415: 57//Z82209

57/IZCZCU9
F-HEMBB1001880//Human genomic DNA sequence from clone 30801 on chromosome Xp11.3-11.4. Contains EST, CA repeat, STS, GSS, CpG Island J/1.0e-18:729:60/IZ93403

F-HEMBB1001899//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from contig 4-10, complete sequence.//0.0038:425:58//AL010216

55 F-HEMBB1001905//S.pombe chromosome III cosmid c330.//1.1e-23:520:62//AL031603

F-HEMBB1001906

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F-HEMBB1001908//Human monocytic leukaemia zinc finger protein (MOZ) mRNA, complete cds.//3.7e-82:672: 81//U47742

- F-HEMBB1001910//Plasmodium falciparum 3D7 chromosome 12 PFYAC181 genomic sequence, WORKING DRAFT SEQUENCE, 8 unordered pieces.//0.0033:566:55//AC005505
- F-HEMBB1001911//Arabidopsis thaliana chromosome II BAC F26C24 genomic sequence, complete sequence.//
  1.0:581:58//AC004705
- F-HEMBB1001915//Caenorhabditis elegans cosmid T05H10, complete sequence.//1.2e-16:283:67//Z47812
   F-HEMBB1001921//Homo sapiens chromosome 17, clone hCIT.123\_J\_14, complete sequence.//3.4e-07.803:58// AC003950
  - F-HEMBB1001922//Plasmodium falciparum chromosome 2, section 28 of 73 of the complete sequence.//5.0e-06: 756:56//4F001391
- F-HEMBB1001925//Human DNA sequence from PAC 212P9 on chromosome 1p34.1-1p35. Contains delta opiate receptor, CpG island, CA repeat, //3.1e-45:609:73//AL009181
  - F-HEMBB1001930//Homo sapiens genomic DNA of 8p21.3-p22 anti-oncogene of hepatocellular colorectal and non-small cell lung cancer, segment 10/11///3/2e-158:745:99//AB020867
- F-HEMBB1001944//, complete sequence.//4.1e-60:638:73//AC005815
- 15 F-HEMBB1001945/HS\_3185\_B1\_G05\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3185 Col=9 Row=N, genomic survey sequence.//1.0:280:58//AQ188882
  - F-HEMBB1001947//Human mRNA for KIAA0392 gene, partial cds.//5.6e-20:333:66//AB002390
    - F-HEMBB1001950//Human lipocortin (LIP) 2 gene, upstream region.//0.0094:180:63//M62899
  - F-HEMBB1001952//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 101A4, WORKING DRAFT SEQUENCE //5 4e-19:329:70//793341
  - F-HEMBB1001953//Homo sapiens chromosome 17, clone hRPK.795\_F\_17, complete sequence.//0.11:589:58// AC005284
    - F-HEMBB1001957//Human DNA sequence from PAC 204E5 on chromosome 12. Contains exon similar to Wilms' Tumour-related protein QM-like P2X-like receptor, ATP ligand gated ion channel, ESTs, CpG island.//9.8e-25:446:
    - F-HEMBB1001962//Homo sapiens chromosome 16, BAC clone 462G18 (LANL), complete sequence.//2.8e-147: 727-97//AC005736
    - F-HEMBB1001967//Homo sapiens clone DJ1102A12, WORKING DRAFT SEQUENCE, 15 unordered pieces.// 3.2e-56:650:71//AC004963
- 30 F-HEMBB1001973//Homo sapiens chromosome 12p13.3-cione RPCI11-350L7, WORKING DRAFT SEQUENCE, 72 unordered pieces //1.2e-42:327:84//AC005844
  - F-HEMBB1001983//CIT-HSP-2315M4.TF CIT-HSP Homo sapiens genomic clone 2315M4, genomic survey sequence.//8.8e-35:198:96//Q028071
    - F-HEMBB1001988//D.polychroa microsatellite sequence (clone Dp 1C e12).//4.5e-07:337:62//X92189
- 39 F-HEMBB1001990/IHS 3234\_A1\_G08\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3234 Col=15 Row=M, genomic survey sequence./i0.039:279:59/IAQ204689 F-HEMBB1001996/Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 191J18, WORKING
  - DRAFT SEQUENCE.//0.18:392:58//AL024507
- F-HEMBB1001997//Homo sapiens clone RG140B11, WORKING DRAFT SEQUENCE, 1 unordered pieces.//1.3e-43:446:71//AC005069
  - F-HEMBB1002002//Plasmodium falciparum 3D7 chromosome 12 PFYAC812 genomic sequence, WORKING DRAFT SEQUENCE, 8 unordered pieces.//0.077:444:58//AC004153
  - F-HEMBB1002005//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 963K23, WORKING DRAFT SEQUENCE.//3.4e-16:173:78//AL031685
- 45 F-HEMBB1002009//Plasmodium falciparum 3D7 chromosome 12 PFYAC357 genomic sequence, WORKING DRAFT SEQUENCE, 7 unordered pieces.//0.00033:790:56//AC005506
  - F-HEMBB1002015//Homo sapiens genomic DNA, chromosome 21q11.1, segment 27/28, WORKING DRAFT SE-QUENCE.//6.7e-05:126:76//AP000056
  - F-HEMBB1002042//Oncorhynchus mykiss cytochrome P450 (CYP4V1) mRNA, partial cds.//6.4e-33:402:69// AF046012
  - F-HEMBB1002043

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- F-HEMBB1002044//Homo sapiens chromosome 5p, BAC clone 50g21 (LBNL H154), complete sequence.//3.0e-167:809:97//AC005740
- F-HEMBB1002045
- F-HEMBB1002049//Homo sapiens chromosome 17, clone hRPC.161\_P\_9, complete sequence.//0.87:177:65//
  AC006237
  - F-HEMBB1002050//Streptomyces coelicolor cosmid D78.//8.5e-08:644:58//AL034355
  - F-HEMBB1002068//Homo sapiens mRNA for KIAA0612 protein, partial cds.//2.5e-05:402:61//AB014512

F-HEMBB1002069

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- F-HEMBB1002092//\*\*\* SEQUENCING IN PROGRESS \*\*\* Homo sapiens chromosome 4, BAC clone B33108; HTGS phase 1, WORKING DRAFT SEQUENCE, 10 unordered pieces.//7.8e-104:550:83//AC004064
- F-HEMBB1002094//Homo sapiens genomic DNA, 21q region, clone: 125H6N2, genomic survey sequence.//2.9e-49:302:83//AG001476
  - F-HEMBB1002115//Homo sapiens chromosome 16, cosmid clone 378E2 (LANL), complete sequence.//0.00023: 542:61//AC004035
- F-HEMBB1002134//Human h-neuro-d4 protein mRNA, complete cds.//7.3e-43:533.70//J43843
  F-HEMBB1002139//HS-1048-A2-B02-MR.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone
  Plate=CT 831 Col=4 Row=C, genomic survey sequence://i0.055/228/66//B38714
  - F-HEMBB1002142//Plasmodium falciparum DNA\*\*\* SEQUENCING IN PROGRESS\*\*\* from MAL1P5, WORKING DRAFT SEQUENCE //0.0095/276/64//AL031748
  - F-HEMBB1002152//Human Chromosome X, WORKING DRAFT SEQUENCE, 4 unordered pieces.//0.055:520: 57//AC002421
- 15 F-HEMBB1002189//Homo sapiens cosmid ICRFc104I0935Q8 from Xq28, complete sequence.//2.6e-05:311:63// AF002998
  - F-HEMBB1002190//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-420 genomic sequence, WORKING DRAFT SEQUENCE, 14 unordered pieces.//5.4e-05:647:59//AC005140
  - F-HEMBB1002193//Sequence 5 from patent US 5709858.//1.8e-34:179:100///80846
- 20 F-HEMBB1002217//Homo sapiens mRNA for zinc finger protein 10.//1.2e-23:405:67//X52332
  - F-HEMBB1002218/HS\_2056\_B1\_C09\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2056 Col=17 Row=F, genomic survey sequence.//3.3e-45:245:97/IA/Q244711
    - F-HEMBB1002232//Human chromosome 11 72g7 cosmid, complete sequence.//1.9e-21:314:70//U73648 F-HEMBB1002247
- F-H-EMBE1002249/i/homo sapiens DNA sequence from BAC 34l8 on chromosome 6p21.3-22.1. Contains ZNF 134 gene coding for Kruppel related Zinc Finger protein 184, a hnRNP core protein A1 (mouse Fil-2, rat helix destabilizing protein, mouse Topoisomenase-inhibitor suppressed gene TiS) LIKE pseudogene, a HB15 (CD83 antigen precursor) LIKE pseudogene, Ser-tRNA, Glu-tRNA and Met-tRNA (Met-tRNA-) gene 1) genes. Contains ESTs, STSs and GSSs. complete sequence (41, 41-e4-523728)ALQ21918
- 39 F-HEMBB1002254//Human chromosome 16 BAC clone LANL cosmid-440E5, WORKING DRAFT SEQUENCE, 2 unordered pieces.//9.8e-40:315:82//AC002506
  - F-HEMBB1002255//Plasmodium falciparum MAL3P3, complete sequence.//0.0035:312:62//Z98547
  - F-HEMBB1002266//Plasmodium falciparum 3D7 chromosome 12 PFYAC1383 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered pieces.//0.013:469:59//AC005504
- 35 F-HEMBB1002280//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-259H10, complete sequence.//5.3e-18:527:61//AC004682
  - F-HEMBB1002300//Human Chromosome 11 Cosmid cSRL30h11, complete sequence.//8.6e-139:818:88// LI73642
- F-HEMBB1002306//HS\_3109\_A2\_H01\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3109 Col=2 Row=O, genomic survey sequence.//1.3e-75:371:98//AQ148164
- F-HEMBB1002327//HS\_3235\_B2\_G10\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3235 Col=20 Row=N, genomic survey sequence.//3.3e-83:418:97//AQ209752
  - F-HEMBB1002329//CITBI-E1-2503J7.TR CITBI-E1 Homo sapiens genomic clone 2503J7, genomic survey sequence.//3.3e-31:220:88//AQ263402
- 45 F-HEMBB1002340
  - F-HEMBB1002342//Homo sapiens mRNA for putative thioredoxin-like protein.//4.1e-154:724:98//AJ010841 F-HEMBB1002358//Human thymidylate kinase (CDC8) mRNA, complete cds.//3.3e-36:192:98//L16991
  - F-HEMBB1002359//Human Rev interacting protein Rip-1 mRNA, complete cds://1.8e-13:96:96//U55766
  - F-HEMBB1002364//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 376D21, WORKING DRAFT SEQUENCE.//7.5e-24:202:71//Z98946
  - F-HEMBB1002371//Plasmodium falciparum 3D7 chromosome 12 PFYAC812 genomic sequence, WORKING DRAFT SEQUENCE, 8 unordered pleces.//1.9e-06:674:56//AC004153
    - F-HEMBB1002381//Homo sapiens chromosome 16, cosmid clone RT163 (LANL), complete sequence.//0.34:238: 61//AC005222
- 55 F-HEMBB1002383

- F-HEMBB1002387//CIT-HSP-2173E20.TR CIT-HSP Homo sapiens genomic clone 2173E20, genomic survey sequence.//5.2e-17:434:66//B91052
  - F-HEMBB1002409//Human DNA sequence from PAC 84F12 on chromosome Xq25-Xq26.3. Contains glypican-3

- precursor (intestinal protein OCI-5) (GTR2-2), ESTs and CA repeat.//1.2e-56:324:88//AL008712
- F-HEMBB1002415//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 364I1, WORKING DRAFT SEQUENCE.//8.9e-35:334:75//AL031319
- F-HEMBB1002425//Chromosome 22q13 BAC Clone CIT987SK-384D8 complete sequence.//1.0e-36:317:76// U62317
  - F-HEMBB1002442//Rattus norvegicus lin-10 protein homolog (lin-10) mRNA, complete cds.//4.3e-88:296:92//
  - F-HEMBB1002453//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 86D1, WORKING DRAFT SEQUENCE //2.7e-43:419:78//AL034349
- F-HEMBB1002457//Homo sapiens clone DJ0982E09, WORKING DRAFT SEQUENCE, 3 unordered pieces.// 1.3e-27:542-68//AC005534
  - F-HEMBB1002458//HS\_3246\_A2\_G05\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3246 Col=10 Row=M, genomic survey sequence.//3.2e-51:257:99//AQ217993
  - F-HEMBB1002477//Human Grb2-associated binder-1 mRNA, complete cds.//1.9e-87:493:92//U43885
- 15 F-HEMBB1002489

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- F-HEMBB1002492//Arabidopsis thaliana BAC T15B16.//0.028:516:57//AF104919
- F-HEMBB1002495//Homo sapiens chromosome 17, clone hRPK.421\_E\_14, complete sequence.//1.1e-16:297: 68//AC006141
- F-HEMBB1002502//Homo sapiens clone DJ1163L11, complete sequence.//1.1e-91:675:82//AC005230
- F-HEMBB1002509//Homo sapiens Chromosome 16 BAC clone CIT987SK-A-248F7, complete sequence.//2.7e-11:648:60//AC004605
  - F-HEMBB1002510//HS\_3236\_B1\_H11\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3236 Col=21 Row=P, genomic survey sequence.//1.2e-06:67:94//AQ205992
  - F-HEMBB1002520//Homo sapiens BAC clone NH0004N07 from Y, complete sequence.//1.2e-70:580:72//
- F-HEMBB1002522//Homo sapiens Xp22 bin 150 clone GSHB-223P11 (Genome Systems Human BAC library) complete sequence.//5.6e-22:516:64//AC004553 F-HEMBB1002531
  - F-HEMBB1002534//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 668J24, WORKING DRAFT SEQUENCE.//6.9e-62:265:87//AL034346
- 30 F-HEMBB1002545//Human BAC clone RG128M16 from 7q21-7q22, complete sequence.//2.7e-44:200:82// AC000059
  - F-HEMBB1002550//Homo sapiens PAC clone DJ0910I17 from 7q11.21-q11.23, complete sequence.//0.22:161: 68//ΔC004927
  - F-HEMBB1002556//Homo sapiens PAC clone DJ0696N01 from 7p21-p22, complete sequence.//7.5e-43:306:77// AC004861
- F-HEMBB1002579
  - F-HEMBB1002582//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 349A12, WORKING DRAFT SEQUENCE.//0.00018:431:61//AL033520
- F-HEMBB1002590//Yeast (S.cerevisiae) mitochondrial apocytochrome b gene, 3' flank.//0.78:147:64//J01471
- 40 F-HEMBB1002596//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from done 9E21, WORKING DRAFT SEQUENCE.//3.6e-50:692:69//AL008639
  - F-HEMBB1002600//Homo sapiens tetraspan NET-5 mRNA, complete cds.//9.1e-151:710:98//AF089749
  - F-HEMBB1002601//Human BAC clone RG020D02 from 7q22, complete sequence.//1.5e-07:416:60//AC002381 F-HEMBB1002603//Human BAC clone GS552A01 from 7q21-q22, complete sequence.//0.40:341:60//AC002454
- 45 F-HEMBB1002607//Mus musculus homeobox containing nuclear transcriptional factor Hmx1 (Hmx1) gene, complete cds://0.0042:460:60//AF009614
  - F-HEMBB1002610//Homo sapiens Chromosome 12q24 PAC RPCI3-462E2 (Roswell Park Cancer Institute Human PAC library) complete sequence.//6.3e-23:559:63//AC003029
  - F-HEMBB1002613//Homo sapiens Chromosome 22q12 BAC Clone 566c1, complete sequence.//4.2e-17:441:63// AC000025
  - F-HEMBB1002614//Plasmodium falciparum chromosome 2, section 54 of 73 of the complete sequence.//0.013: 324:56//AE001417
  - F-HEMBB1002617//Homo sapiens chromosome 16 BAC clone CIT987SK-334D11 complete sequence.//2.1e-07:
- 55 F-HEMBB1002623//C.hvalina microsatellite marker DNA (id ATCC4).//0.57:106:66//Z95304
  - F-HEMBB1002635//Human JNK3 alpha2 protein kinase (JNK3A2) mRNA, complete cds.//4.8e-22:127:100//
    - F-HEMBB1002664//HS 2265 A1 H06 MF CIT Approved Human Genomic Sperm Library D Homo sapiens ge-

- nomic clone Plate=2265 Col=11 Row=O, genomic survey sequence.//0.54:115:67//AQ101557
- F-HEMBB1002677//Homo sapiens (subclone 3\_d1 from P1 H25) DNA sequence, complete sequence.//2.2e-49: 784:68//L81774
- F-HEMBB1002683//Homo sapiens type IV collagen 5a chain (COL4A5) gene, exon 23.//1.0:112:63//U04492
- F-HEMBB1002684//HS-1050-A2-G06-MF.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 772 Col=12 Row=M, genomic survey sequence.//4.4e-07:86:84//B39748
  - F-HEMBB1002686//HS-1023-B2-F10-MR.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 802 Col=20 Row=L, genomic survey seguence //0.98:183:61//B34077
  - F-HEMBB1002692//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 1108H3, WORKING DRAFT SEQUENCE //0.00039:408:60//AL033525
    - F-HEMBB1002697//Homo sapiens clone DJ1087M19, WORKING DRAFT SEQUENCE, 7 unordered pieces.//
      7.3e-35:323:74//AC004955
  - F-HEMBB1002699//Mus musculus D6MM5e protein (D6Mm5e) and DOK protein (Dok) genes, complete cds; and LOR2 protein (Lor2) gene, partial cds.//0.031;325:62//AF084363
- F-HEMBB1002702/iHS-1025-A2-D01-MR.abi CIT Human Genomic Sperm Library C Homo sapiens genomic clone Plate=CT 804 Col=2 Row=G, genomic survey sequence.//1.8e-25:158:95/IB34720
  - F-HEMBB1002705//Homo sapiens DNA, chromosome 21q22.2, PAC clone 25P16 complete sequence, encoding carbonyl reductase and carbonyl reductase 3 (complete cds).//1.7e-137:534.96//AB003151
  - F-HEMBB1002712//Human DNA sequence from cosmid cU115G11, between markers DXS6791 and DXS8038 on chromosome X contains ESTs and STS.//0.0019:612:58//Z71187
  - on chromosome x contains ES is and STS://U.U019:612:58//Z/1187 F-MAMMA1000009//Human chromosome 1 BAC 308G1 genomic sequence, WORKING DRAFT SEQUENCE, 3 unordered pieces.//6.1e-43:354:81/MC003117
  - F-MAMMA1000019

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- F-MAMMA1000020//H.sapiens mRNA for flavin-containing monooxygenase 5 (FMO5),//2.0e-40:185:97//Z47553
  F-MAMMA1000025//Homo sapiens PAC clone DJ0806A17 from 7p13-p14, complete sequence.//1.0:211:65//
  - AC005483
    F-MAMMA1000043//Human angiotensin I-converting enzyme (ACE) gene, intron 12,//0.075;204;65//M73275
  - F-MAMM1000045//Human DNA sequence from clone 142F18 on chromosome Xq26.3-27.2 Contains part of a gene similar to melanoma-associated antigen, EST, GSS and an inverted repeat, complete sequence.//4.1e-122: 485-78/JL 031072
- F-MAMMA1000055//M.musculus mRNA for testin.//2.1e-35:559:66//X78989
  - F-MAMMA1000057//Homo sapiens chromosome 17, clone hRPK.259\_G\_18, complete sequence.//5.5e-121:703: 89//AC005829
  - F-MAMMA1000069//Homo sapiens minisatellite ceb1 repeat region.//0.00013:329:60//AF048727
- F-MAMMA1000084//Homo sapiens DNA sequence from PAC 958B3 on chromosome Xp22.11-Xp22.22. Contains ESTs STS and CpG island.//2.1e-53:445:79//Z93023
  - F-MAMMA1000085//Caenorhabditis elegans cosmid Y23H5A.//0.0017:164:64//AF077541
  - F-MAMMA1000092//Homo sapiens BAC clone GS465N13 from 7p15-p21, complete sequence.//1.2e-70:598:78// AC004744
- 40 F-MAMMA1000103//Homo sapiens chromosome 17, clone hClT.91\_J\_4, complete sequence.//1.1e-156:857:92// AC003976
  - F-MAMMA1000117/IHS\_3223\_B2\_D08\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3223 Col=16 Row=H, genomic survey sequence./I5-4e-100:527:94/IAQ221160 F-MAMMA1000129/Iryanodine receptor./I0.055-492:59/IA20359
- 45 F-MAMMA1000133

F-MAMMA1000163

- F-MAMMA1000134//HS\_3078\_B1\_C02\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3078 Col=3 Row=F, genomic survey sequence.//2.1e-93:462:97//AQ144362
- F-MAMMA1000139//Homo sapiens Xp22 PAC RPCI1-5G11 (from Roswell Park Cancer Center) complete sequence.//3.3e-14:322:65//AC002369
- F-MAMMA1000143//Homo sapiens mRNA for KIAA0685 protein, complete cds.//6.9e-25.148:97//AB014585 F-MAMMA1000155//Homo sapiens homeobox transcription factor barx2 (BARX2) mRNA, complete cds.//1.0e-29: 219.97//AF031924
  - F-MAMMA1000171//Homo sapiens chromosome 19, CIT-HSP BAC 470n8, complete sequence.//6.3e-14:92:88//
- F-MAMMA1000173//Mus musculus SH3-containing protein SH3P7 mRNA, complete cds. similar to Human Drebrin.//2.2e-114:698:87//U58884
  - F-MAMMA1000175//HS 3050 B1 B03 MF CIT Approved Human Genomic Sperm Library D Homo sapiens ge-

nomic clone Plate=3050 Col=5 Row=D, genomic survey sequence.//6.2e-73:357:99//AQ102678

F-MAMMA1000183//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone Y313F4, WORKING DRAFT SEQUENCE.//4.6e-94:904:73//AL023808

F-MAMMA1000198//Z.diploperennis repetitive DNA (clone ZEAR 266).//0.18:152:70//X53610

F-MAMMA1000221//Human Chromosome 15q11-q13 PAC clone pDJ778a2, complete sequence.//0.017:99:75//

F-MAMMA1000227//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 467K16, WORKING DRAFT SEQUENCE.//0.36:312:62//AL031283

F-MAMMA1000241//HS\_3217\_B1\_B02\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3217 Col=3 Row=D, genomic survey sequence.//1.9e-94:456:98//AQ193401

F-MAMMA1000251//Homo sapiens NF2 gene.//0.00092:270:64//Y18000

F-MAMMA1000254//Plasmodium falciparum 3D7 chromosome 12 PFYACB8-420 genomic sequence, WORKING DRAFT SEQUENCE, 14 unordered pieces //0.0034;777:57//AC005140

F-MAMMA1000257//Homo sapiens DNA sequence from PAC 201D7 on chromosome 6p22.1-22.3. Contains EST and STS://0.00036:230:65//AL022717

F-MAMMA1000264//Homo sapiens (subclone 9\_f5 from P1 H17) DNA sequence, complete sequence.//1.5e-30:

F-MAMMA1000266//Bacillus lynceorum strain pMEL12 Bag320 satellite DNA.//0.28;218;64//AF034430

F-MAMMA1000270//Human Chromosome 16 BAC clone CIT987SK-A-270G1, complete sequence.//1 .4e-157: 788:96//AF001549

F-MAMMA1000277//Mycobacterium tuberculosis H37Rv complete genome; segment 48/162.//0.70:320:61// AL021897

F-MAMMA1000278//Sequence 23 from patent US 5708157.//9.3e-103:540:95///80055

F-MAMMA1000279//Human DNA sequence from clone 769D20 on chromosome Xp21.1-21.3 Contains EST, STS, GSS, complete sequence //2.4e-49:262:77//AL031643

F-MAMMA1000284/lcSRL-165E12-u cSRL flow sorted Chromosome 11 specific cosmid Homo sapiens genomic clone cSRL-165E12, genomic survey sequence.//1.1e-30:324:75//B03004

F-MAMMA1000287//Homo sapiens, clone hRPK.15\_A\_1, complete sequence.//2.7e-54:401:83//AC006213 F-MAMMA1000302//Drosophila melanogaster complete mitochondrial genome.//0.0051:307:61//U37541

30 F-MAMMA100307//Homo sapiens chromosome 12p13.3 clone RPCI5-1154L15, WORKING DRAFT SE-QUENCE 67 upordered pieces //0.15/449/59//AC006205

F-MAMMA1000309//cDNA coding human apolipoprotein E3.//0.00010:691:58//E00359

F-MAMMA1000312//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 798A17, WORKING DRAFT SEQUENCE //0.27:301:60//AL031274

F-MAMMA1000313 F-MAMMA1000331

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F-MAMMA1000331//Human Chromosome 16 BAC clone CIT987SK-A-735G6, complete sequence.//9.8e-06:151: 71//AC002400

F-MAMMA1000339

F-MAMMA1000340//HS\_2181\_B2\_F07\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2181 Col=14 Row=L, genomic survey sequence.//4.3e-05:181:68//AQ024288

F-MAMMA1000348//Homo sapiens chromosome 17, clone HRPC843B9, complete sequence.//5.3e-30:575:66// AC004139

F-MAMMA1000356//Homo sapiens clone RG038K21, WORKING DRAFT SEQUENCE, 3 unordered pieces.// 1.8e-52:264:76//AC005052

45 F-MAMMA1000360//Homo sapiens PAC clone DJ0755G17 from 7p21-p22, complete sequence.//6.5e-91:569:88// AC004879

F-MAMMA1000381//Human DNA sequence from PAC 507115 on chromosome Xq26.3-27.3. Contains 60S ribosonal protein L44 (L41, L36) like gene, ESTs, STSs and a polymorphic CA repeat./1.4.e-42.315:83/Z98950 F-MAMMA1000372//Human DNA sequence ""SEQUENCING IN PROGRESS "" from clone Y738F9, WORKING

DRAFT SEQUENCE.//Z.9e-114:516:89//AL022345
F-MAMMA1000385//CITBI-E1-2517E13.TF CITBI-E1 Homo sapiens genomic clone 2517E13, genomic survey

sequence./I/6.9e-26:377:71//AQ279944
F-MAMA100388/Homo sapiens UKLF mRNA for ubiquitous Kruppel like factor, complete cds.//3.7e-148:710:
98/JAB015132

55 F-MAMMA1000395

F-MAMMA1000402//Homo sapiens clone DJ0718N17, complete sequence.//4.0e-115:845:85//AC005999

F-MAMMA1000410//HS\_3245\_A1\_C02\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3245 Col=3 Row=E, genomic survey sequence.//9.6e-42:350:80//AQ205768

- F-MAMMA1000413//HS\_3223\_B2\_F01\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3223 Col=2 Row=L, genomic survey sequence.//1.6e-48:318:89//AQ188456
- F-MAMMA1000414//HS\_2027\_B2\_C04\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2027 Col=8 Row=F, genomic survey sequence.//1.4e-46:286:92//AQ231369
- F-MAMMA1000416//Drosophila melanogaster DNA sequence (P1s DS07528 (D169) and DS06665 (D220)), complete sequence.//9.4e-33:310:72//AC004640
  - F-MAMMA1000421//Homo sapiens clone DJ1129D05, complete sequence.//3.3e-29:223:84//AC005630 F-MAMMA1000422
  - F-MAMMA1000423//Drosophila vakuba mitochondrial DNA molecule.//2.2e-10:639:57//X03240
- F-MAMMA1000424//Homo sapiens DNA from chromosome 19, BAC 33152, complete sequence.//4.6e-47:556: 68//AC003973
  - F-MAMMA1000429//Mus musculus SDP8 mRNA, complete cds.//8.0e-99:545:92//AF062484

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- F-MAMMA1000431//Homo sapiens clone DJ1039L24, WORKING DRAFT SEQUENCE, 3 unordered pieces.//
  4.8e-41:289:79//AC005283
- F-MAMMA1000444//Human DNA sequence from clone 714B7 on chromosome 22q12.2-13.2 Contains CYTO-CHROME C OXIDASE VIIB precursor like pseudogene and ESTs, complete sequence./I/2.3e-34:291:80/I/299755 F-MAMMA1000148
  - F-MAMMA1000458//Mus musculus clone OST9003, genomic survey seguence.//5.0e-53:231:84//AF046620
  - F-MAMMA1000468//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 291J10, WORKING DRAFT SEQUENCE.//0.75;303:60//Z93017
  - F-MAMMA1000472//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 414D7, WORKING DRAFT SEQUENCE.//4.0e-41:403:77//AL033543
    - F-MAMMA1000478//Homo sapiens clone RG270D13, WORKING DRAFT SEQUENCE, 18 unordered pieces.// 9.5e-54:369:77//AC005081
- F-MAMMA1000483//Homo sapiens Chromosome 16 BAC clone CIT987SK-44M2, complete sequence.//3.6e-34: 332-77//AC004381
  - F-MAMMA1000490//Homo sapiens 12q13.1 PAC RPCI1-90J4 (Roswell Park Cancer Institute Human PAC library) complete sequence.//8.9e-128:822:87//AC003686
- F-MAMMA1000500//CIT-HSP-231905.TF CIT-HSP Homo sapiens genomic clone 231905, genomic survey sequence.//4.8e-29:175:94//AQ044812
- F-MAMMA1000501//Homo sapiens DNA sequence from clone 78F24 on chromosome 22q12.1-12.3. Contains one exon of an Oxysterol-binding protein (OSBP) LIKE gene. Contains GSSs and an STS, complete sequence.// 5.79-4-5334.82/IAI.023336
- F-MAMMA1000516//Human DNA sequence from PAC 50A13 on chromosome Xp11. Contains ATP SYNTHASE LIPID BINDING PROTEIN P1 (P2, P3) precursor (ATPSG1, ATPSG2, ATPSG3) like pseudogene, ESTs and STSs. Contains polymorphic CA repeat.//2.9e-43:529:69/J292545
  - F-MAMMA1000522//Human DNA sequence from clone 20J23 on chromosome Xq26.2-27.2 Contains ras-related C3 bublium toxin substrate 1 (22-1RAC1) (ras-like protein TC25) EST, CA repeat, STS, CpG island, complete sequence //2.0e-14:380-63//AL022576
- 40 F-MAMMA1000524//Homo sapiens chromosome 10 clone CIT-HSP-1338F24 map 10p11.2-10p12.1, complete sequence.//1.4e-22:420:66//AC006101
  - F-MAMMA1000559//Human HepG2 3' region cDNA, clone hmd3f08,//5.4e-29:168:97//D16922
  - F-MAMMA1000565//RPCI11-61K6.TJ RPCI11 Homo sapiens genomic clone R-61K6, genomic survey sequence.//
    1.7e-120:561:100//AQ194238
- 45 F-MAMMA1000567//Human DNA sequence from PAC 179D3, between markers DXS6791 and DXS8038 on chromosome X contains S10 GTP-binding protein, ESTs and CpG island.//3.1e-43:387:80//Z81370
  - F-MAMMA1000576//Homo sapiens BAC clone RG442F18 from 2, complete sequence.//1.2e-30:237:75// AC005104
  - F-MAMMA1000583//RPCI11-60M22.TJ RPCI11 Homo sapiens genomic clone R-60M22, genomic survey sequence.//9.6e-102:487:99//AQ198091
  - F-MAMMA1000585//Homo sapiens clone UWGC:djs14 from 7p14-15, complete sequence://5.2e-39:370:78// AC006195
  - F-MAMMA1000594//Homo sapiens chromosome 19, cosmid R31646, complete sequence.//3.9e-43:328:83// AC005338
- F-MAMMA1000597//Homo sapiens chromosome 17, clone hRPK.481\_C\_4, complete sequence.//1.5e-32:259: 82//AC005839
  - F-MAMMA1000605//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 97P20, WORKING DRAFT SEQUENCE //2.4e-59:318:83//AL031297

- F-MAMMA1000612//HS\_2188\_A2\_D02\_MF CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2188 Col=4 Row=G, genomic survey sequence.//4.8e-30:171:96//AQ116793
- F-MAMMA1000616//HS\_3176\_A1\_E06\_T7 CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3176 Col=11 Row=I, genomic survey sequence.//4.7e-28:287:79//AQ300310
- F-MAMMA1000621//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from done 273F20, WORKING DRAFT SEQUENCE.//0.015:478:58//AL034371
  - F-MAMMA1000623

- F-MAMMA1000625//DNA encoding Hepatitis C virus antigen.//0.93:196:61//E06898
- F-MAMMA1000643//Homo sapiens nephrocystin (NPHP1) mRNA, partial cds.//0.95:365:59//AF023674
- F-MAMMA1000664//HS\_3096\_B1\_C02\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=3096 Col-3 RowerF, genomic survey sequence./IZ.7e-51:257:99/IA5137
  F-MAMMA1000669//Homo sapiens chromosome 19, cosmid R56098. compile sequence./IZ.0e-66:586:67//
  - F-MAMMA1000669//Homo sapiens chromosome 19, cosmid R26908, complete sequence.//2.0e-66:586:67/ AC004785
- F-MAMMA1000670//HS\_2243\_B2\_A08\_MR CIT Approved Human Genomic Sperm Library D Homo sapiens genomic clone Plate=2243 Col=16 Row=B, genomic survey sequence.//8.7e-05:94:80//AQ153650
- F-MAMMA1000672//Mus musculus clone OST8270, genomic survey sequence.//3.9e-64:471:81//AF046705
  - F-MAMMA1000684//Suid herpesvirus 1 Rsp40 mRNA, partial cds.//1.2e-07:1866/7/U27489
    F-MAMMA1000696//Human oligodendrocyte myelin glycoprotein (OMG) exons 1-2; neurofibromatosis 1 (NF1) exons 28-49; ecotropic viral integration site 28 (EVI2B) exons 1-2; ecotropic viral integration site 2A (EVI2A) exons
- 1-2; adenylate kinase (AK3) exons 1-2.//3.0e-53:653:70//L05367 F-MAMMA1000707//CIT-HSP-2302019.TR CIT-HSP Homo sapiens genomic clone 2302O19, genomic survey se
  - quence.//1.8e-08:131:77//AQ017947
    F-MAMMA1000713//Rattus norvegicus clonel polymeric immunoglobulin receptor mRNA 3' untranslated region,
- GA rich region, and microsatellites with GGA-triplet and GAA-triplet repeats.//0.062:134:67//U00762

  5 F-MAMMA1000714//Chicken hsp90 gene for 90 kDa-heat shock protein 5-end.//1.0:266:61//X15028
  - F-MAMMA1000718//CIT-HSP-2171B10.TF CIT-HSP Homo sapiens genomic clone 2171B10, genomic survey sequence.//3.6e-05;289:60//B95401
  - F-MAMMA1000720//Homo sapiens chromosome 19, cosmid R33632, complete sequence.//4.4e-184:842:98// AC005781
- 30 F-MAMMA1000723//Homo sapiens clone DJ0892G19, complete sequence.//8.8e-05:430:60//AC004917
  - F-MAMMA1000731//Drosophila melanogaster DNA sequence (P1 DS07049 (D133)), complete sequence.//3.8e-55/796.66/IAC004274
    F-MAMMA1000732//Homo sapiens chromosome 21a22.3 PAC 141B3. complete sequence. containing ribosomal
    - protein homologue pseudogene L23a.//6.6e-77:555:74//AF064859
- 35 F-MAMMA1000733//Plasmodium falciparum DNA \*\*\* SEQUENCING IN PROGRESS \*\*\* from MAL1P6, WORK-ING DRAFT SEQUENCE.//0.98:479:58//AL031749
  - F-MAMMA1000734//Homo sapiens SEC63 (SEC63) mRNA, complete cds.//7.3e-168:802:98//AF100141
  - F-MAMMA1000738//S.cerevisiae chromosome XIV reading frame ORF YNL132w.//8.6e-31:626:63//Z71408
    F-MAMMA1000744//Gorilla Alu-repetitive sequence in beta-globin gene cluster.//2.7e-54:410:82//X06123
- 40 F-MAMMA1000746/Homo sapiens Chromosome 16 BAC clone CIT987SK-A-10F4, complete sequence.//3.7e-109:779:83//AC004158
  - F-MAMMA1000752//Homo sapiens clone RG219E16, WORKING DRAFT SEQUENCE, 3 unordered pieces.// 1.2e-20:444:63//AC005075
  - F-MAMMA1000760//Homo sapiens clone RG015P03, complete sequence.//1.5e-44:403:79//AC005048
- 45 F-MAMMA1000761//Homo sapiens Chromosome 7 BAC Clone 239c10, WORKING DRAFT SEQUENCE, 9 unordered pieces.//2.3e-22:159:81//AC004166
  - F-MAMMA1000775//Homo sapiens chromosome 17, clone hRPK.849\_N\_15, complete sequence.//1.3e-51:789: 68//AC005703
  - F-MAMMA1000776//Human DNA sequence from BAC 57G9 on chromosome 22q12.1 Contains ESTs, CA repeat, GSS.//5.7e-40:238:78//Z95116
    - F-MAMMA1000778//Human DNA sequence \*\*\* SEQUENCING IN PROGRESS \*\*\* from clone 153G14, WORKING DRAFT SEQUENCE //7.6e-29:222:84//AL031118
  - F-MAMMA1000782//Human 2.4-dienoyl-CoA reductase gene, exon 9.//0.90:137:62//U94987
- F-MAMMA1000798//\*\* SEQUENCING IN PROGRESS \*\*\* EPMI/APECED region of chromosome 21, clones A68E8, B127P21, B1731.3, B23N8, C1242C9, C579E2, A7086, B15909, B175D10, B52C10, C12461 Note: Sequencing in this region has been discontinued by the Stanford Human Genome Center, WORKING DRAFT SEQUENCE. 50 unordered pieces./N, 000581:835.71/Mc003656
  - F-MAMMA1000802//Homo sapiens chromosome 19, cosmid R33729, complete sequence.//6.3e-151:714:99//